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4 **Abscisic Acid-Triggered Persulfidation of Cysteine Protease ATG4
5 Mediates Regulation of Autophagy by Sulfide**

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23 Short title: ATG4 is a target of sulfide

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31 **ABSTRACT**

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33 Hydrogen sulfide is a signaling molecule that regulates essential processes in plants,
34 such as autophagy. In *Arabidopsis thaliana*, hydrogen sulfide negatively regulates
35 autophagy independently of reactive oxygen species via an unknown mechanism.
36 Comparative and quantitative proteomic analysis was used to detect abscisic acid-
37 triggered persulfidation that reveals a main role in the control of autophagy mediated by
38 cysteine protease AtATG4a. This protease undergoes specific persulfidation of Cys170
39 that is a part of the characteristic catalytic Cys-His-Asp triad of cysteine proteases.
40 Regulation of the ATG4 activity by persulfidation was tested in a heterologous assay
41 using the *Chlamydomonas reinhardtii* CrATG8 protein as a substrate. Sulfide
42 significantly and reversibly inactivates AtATG4a. The biological significance of the
43 reversible inhibition of the ATG4 by sulfide is supported by the results obtained in
44 Arabidopsis leaves under basal and autophagy-activating conditions. A significant
45 increase in the overall ATG4 proteolytic activity in Arabidopsis was detected under
46 nitrogen starvation and osmotic stress and can be inhibited by sulfide. Therefore, the
47 data strongly suggest that the negative regulation of autophagy by sulfide is mediated by
48 specific persulfidation of the ATG4 protease.

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53 **INTRODUCTION**

54

55 Hydrogen sulfide is currently recognized as a signaling molecule. In plant systems, H₂S
56 is considered to be as important as NO and H₂O₂ and regulates essential processes of
57 plant performance (Garcia-Mata and Lamattina, 2013; Calderwood and Kopriva, 2014;
58 Jin and Pei, 2015; Gotor et al., 2017). Sulfide mediates tolerance against a range of
59 plant stresses from heavy metal toxicity to salinity and drought to enhance plant
60 viability (Gotor et al., 2019). Sulfide regulates critical processes, including autophagy
61 (Alvarez et al., 2012; Gotor et al., 2013; Gotor et al., 2015; Laureano-Marin et al.,
62 2016b; Laureano-Marin et al., 2016a) and the abscisic acid (ABA)-dependent stomatal
63 movement (Jin et al., 2013; Scuffi et al., 2014; Papanatsiou et al., 2015; Scuffi et al.,
64 2018).

65 Despite increasing evidence of the biological function of H₂S, there is a considerable
66 lack of information on the mechanism of action of H₂S in particular physiological
67 processes. The mechanism of action must be related to chemical reactivity of H₂S with
68 other molecules. H₂S was suggested to coordinate the metal centers of metalloproteins
69 (Vitvitsky et al., 2018) or act as a reductant of reactive oxygen species (Zaffagnini et al.,
70 2019). A third mechanism of action of H₂S is based on its ability to modify the thiol
71 group (-SH) of the cysteine residues in target proteins to form a persulfide group (-SSH)
72 resulting in functional changes in the protein structure, activity, or subcellular
73 localizations (Aroca et al., 2015; Aroca et al., 2017b). This posttranslational
74 modification is called persulfidation (known previously as S-sulphydratation) and has
75 been initially demonstrated in the mammalian (Mustafa et al., 2009; Paul and Snyder,
76 2012) and plant systems (Aroca et al., 2015; Aroca et al., 2017a) using specific labeling
77 methods.

78 Our previous investigations in *Arabidopsis thaliana* demonstrated that hydrogen
79 sulfide functions as a signaling molecule in the cytosol that negatively regulates
80 autophagy (Alvarez et al., 2012; Gotor et al., 2013; Romero et al., 2013), which is a
81 highly conserved process involving digestion of the cell contents for recycling and
82 maintenance of cell homeostasis. Autophagy occurs at the basal levels in eukaryotic
83 cells and is induced by internal or external perturbations. In plants, autophagy is
84 involved in development, immune response, and senescence and is induced by stress
85 conditions, including nutrient limitation and other abiotic stresses (Liu and Bassham,
86 2012; Pérez-Pérez et al., 2012; Masclaux-Daubresse et al., 2017; Ustun et al., 2017;

87 Marshall and Vierstra, 2018a). This catabolic process is characterized by *de novo*
88 synthesis of autophagosomes in the cytosol, in which cytoplasmic materials to be
89 recycled are sequestered, transported, and released into the plant vacuole. Various
90 receptors have been described to assist with specific cargo recruitment and degradation
91 via selective autophagy. The core autophagy machinery is highly conserved in all
92 studied eukaryotes, and involved proteins are referred to as autophagy-related (ATG).
93 These ATG proteins include the ATG8 and ATG12 ubiquitin-like conjugation systems
94 that catalyze the covalent attachment of ATG8 to the phospholipid
95 phosphatidylethanolamine (PE), which is an essential adduct for the formation of
96 autophagosomes (Mizushima et al., 2011). Before this conjugation, the C-terminal
97 extension of newly synthesized ATG8 has to be cleaved by the Cys-type protease ATG4
98 to expose a highly conserved Gly, which is necessary for conjugation to PE.
99 Additionally, ATG4 functions as a deconjugating enzyme that cleaves the amide bond
100 between ATG8 and PE allowing the recycling of free ATG8 (Nair et al., 2012;
101 Nakatogawa et al., 2012; Yu et al., 2012).

102 The role of sulfide as a repressor of autophagy is independent of nutrient conditions
103 and specific tissues because sulfide inhibits autophagy in leaves under dark-induced
104 carbon starvation (Alvarez et al., 2012) or in roots under nitrogen deprivation
105 (Laureano-Marin et al., 2016a), and both conditions are unrelated to sulfur metabolism.
106 A study aiming to decipher the mechanism of action of H₂S showed that it is
107 independent of the formation of reactive oxygen species (ROS), such as hydrogen
108 peroxide or superoxide anions, and therefore H₂S does not serve as a reducer in the
109 regulation of autophagy (Laureano-Marin et al., 2016a). Interestingly, a comparative
110 and quantitative proteomic analysis was performed to detect endogenous persulfidated
111 proteins; the results indicated that at least 10% of the entire *Arabidopsis* proteome
112 undergoes persulfidation under physiological conditions suggesting a widespread
113 distribution of this posttranslational modification (Aroca et al., 2017a). Furthermore,
114 persulfidation of various components of the ABA signaling pathway has been recently
115 described as a specific mechanism of action by which H₂S controls the guard cell ABA
116 signaling (Chen et al., 2020; Shen et al., 2020).

117 In this study, a comparative and quantitative proteomic analysis was used to detect
118 persulfidated proteins in the leaves of *Arabidopsis* exogenously treated with ABA.
119 Interestingly, comparison of the untreated and ABA-treated samples indicated that
120 AtATG4a was the protein with the highest difference in the persulfidation level. We

121 then sought to determine whether persulfidation is the mechanism of the regulation of
122 autophagy by sulfide in Arabidopsis under the stress conditions and to ascertain whether
123 persulfidation regulates the activity of ATG4. To this aim, an enzymatic assay using *C.*
124 *reinhardtii* CrATG8 as ATG4 substrate was developed. In-depth analysis of ATG4
125 proteolytic activity was performed *in vitro* and in a cell-free system using total extracts
126 from Arabidopsis under basal or autophagy-activating conditions, including nitrogen
127 limitation or osmotic stress. The results indicate that ATG4 is a specific target of
128 persulfidation.

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130

132 **RESULTS**

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134 **Autophagy induction by abscisic acid-triggered persulfide modification is
135 mediated by AtATG4a regulation by hydrogen sulfide in Arabidopsis**

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137 Abscisic acid can trigger changes in hydrogen sulfide level and protein persulfide
138 modification in the guard cells to modulate stomatal closure (Shen et al., 2020). To
139 assess whether ABA also regulates protein persulfidation levels in the mesophyll cells, a
140 sequential window acquisition of all theoretical spectra-mass spectrometry (SWATH-
141 MS) quantitative approach was combined with the tag-switch method to measure
142 protein persulfidation (Aroca et al., 2017a). Protein samples from three biological
143 replicates (independent pools) of leaf tissue treated with ABA for 0 h (control sample),
144 3 h, and 6 h were isolated and subjected to the tag-switch procedure (Figure 1A). The
145 proteins eluted from the streptavidin beads were digested, and the peptide solutions
146 analyzed in two sequential steps: a shotgun data-dependent acquisition (DDA) approach
147 to generate the spectral library and SWATH acquisition by the data-independent
148 acquisition (DIA) method. In the first step, integration of the nine datasets resulted in
149 identification of a total of 10,329 peptides (1% FDR and 90% confidence) and 1,434
150 unique proteins (1% FDR) that were used as a spectral library (Supplemental Dataset 1).
151 In the second step, to quantify protein levels using SWATH acquisition, the same six
152 biological samples were analyzed in two technical replicates by the DIA method using
153 the LC gradient and LC-MS equipment employed in generation of the spectral library.

154 Therefore, six datasets were generated from the control and ABA-treated (for 3 and 6
155 h) samples to yield a total of 18 datasets used for the quantitative analysis. The fragment
156 spectra were extracted for the eighteen runs, and 33,887 ion transitions, 4,871 peptides,
157 and 1,157 proteins were quantified. Principal component analysis (PCA) of the protein
158 sample subgroups revealed significant reproducible data between the replicates and
159 differences between the ABA treatments (Figure 1B).

160 Comparison between the control (0 h) and the 3 h ABA treatment samples (0 h vs 3 h
161 ABA; Supplemental Dataset 2) showed that 192 proteins were more abundant in the
162 control samples with the fold change > 1.5 (p value < 0.05), and 242 proteins were less
163 abundant with the fold change < 0.66 (p value < 0.05) (Supplemental Dataset 3) (Figure
164 1C). Higher abundance of a protein in the control samples than that in the samples
165 prepared after 3 h ABA treatment means that the protein is more persulfidated in the

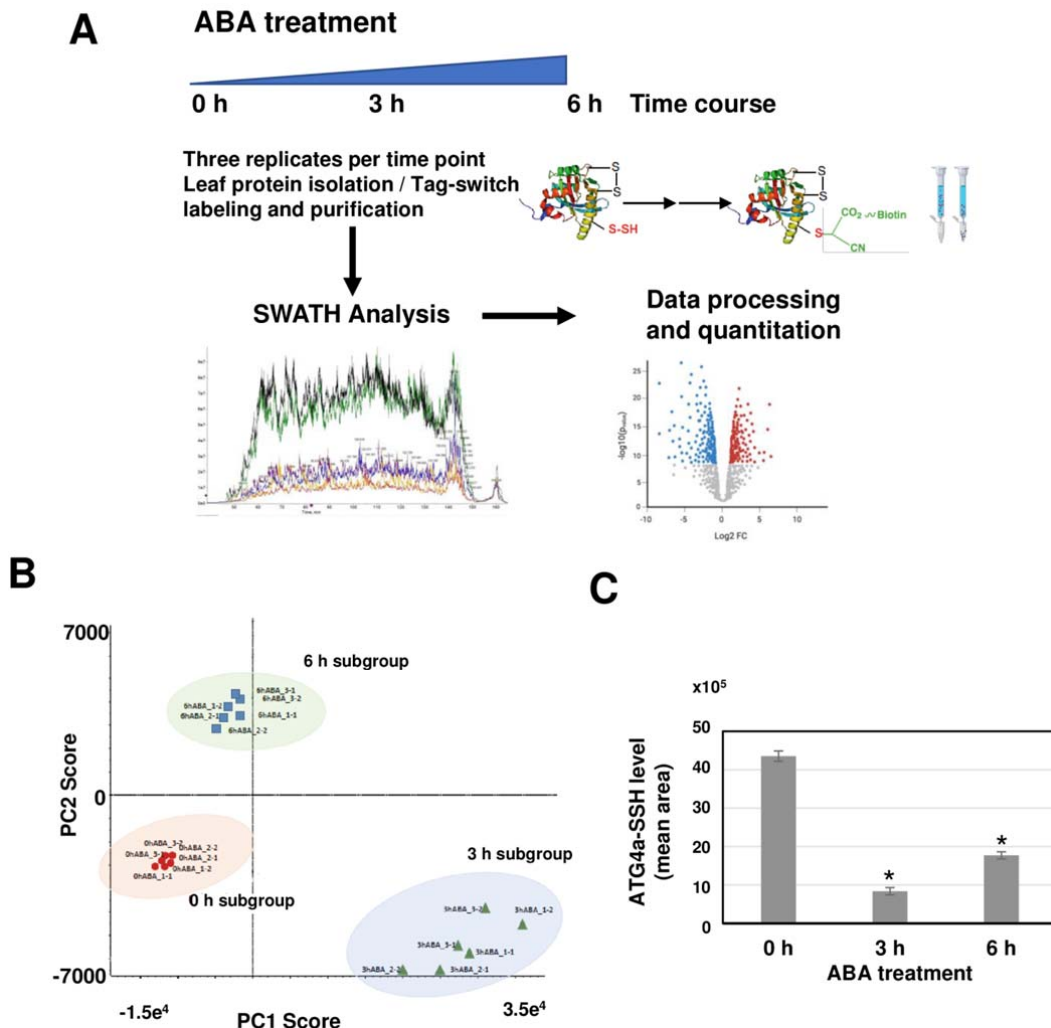


Figure 1. Proteomic analysis of protein persulfidation in response to ABA in mesophyll cells.
(A) Workflow of ABA leaf treatments followed by tag-switch protein labeling with CN-biotin, protein purification, and quantitative SWATH analysis of eluted proteins.
(B) PCA representation plot of the 18 samples after SWATH analysis. Score for PC1 (70 %) versus PC2 (9.9%), Pareto scaling.
(C) Level of persulfidated ATG4a after ABA treatments. Values represent the mean peak areas of extracted ion chromatogram of identified ATG4a peptides. *, p<0.05

166 control because the tag-switch labeling recovers more protein from the streptavidin
 167 column. Lower abundance in the control means that a protein is more persulfidated in
 168 the 3 h ABA samples.

169 A total of 192 proteins with reduced persulfidation level after 3 h of ABA treatment
 170 were analyzed based on their assigned functions and classified into 32 functional groups
 171 using the MapMan nomenclature (Thimm et al., 2004; Klie and Nikoloski, 2012)
 172 (Supplemental Table 1). The most numerous set corresponded to the general protein

173 group (bin 29) (Supplemental Table 2), which included 18.2% of the total identified
174 proteins with 35 total elements, 29 of which are involved in protein amino acid
175 activation (8 elements, tRNA ligases), protein synthesis (12 elements) and protein
176 degradation (9 elements). The latter group included the Cys-type protease AtATG4a
177 involved in autophagy, which showed the highest persulfidation change (5.20-fold
178 change), and the S1 RNA-BINDING RIBOSOMAL PROTEIN 1 (3.69-fold change)
179 that regulates seedling growth in the presence ABA or under abiotic stress conditions
180 (Gu et al., 2015). Additional proteins involved in the regulatory process and with a
181 reduction in their persulfidation levels included MAPK kinase 6 (MPK6, AT2G43790),
182 tyrosine phosphatase 1 (PTP1, AT1G71860), glycine-rich protein 8 (GRP8,
183 AT4G39260), and 9-cis-epoxycarotenoid dioxygenase 4 (NCED4, AT4G19170).

184 The quantitative analysis of the control and the 6 h ABA treatment samples
185 (Supplemental Dataset 4) showed a reduction in the number of regulated proteins; only
186 120 proteins were more abundant in the control *versus* the ABA treatment samples with
187 the fold change > 1.5 (*p* value < 0.05), and 198 proteins were less abundant with the
188 fold change < 0.66 (*p* value < 0.05) (Supplemental Dataset 5) (Figure 1C).

189 Comparison of differentially regulated proteins at 3 and 6 h of ABA treatment
190 showed that 42% of the proteins are regulated under both conditions and had different
191 level of persulfidation. The cysteine protease AtATG4a showed a reduction in the level
192 of persulfidation down to only 2.45-fold after 6 h of ABA treatment compared to that
193 under the control conditions; this level was higher than that in the 3 h ABA treatment
194 samples (Figure 1D). Therefore, persulfidation level of AtATG4a was transiently
195 reduced after a short ABA treatment, and this reduction was very significant compared
196 to the untreated control.

197 The proteomics data suggest that AtATG4a protease is differentially persulfidated
198 depending on the treatment conditions, which may be related with the progress of
199 autophagy. To prove this assumption, the regulation of autophagy by ABA treatments
200 and the effects of sulfide under these conditions were analyzed. Arabidopsis seedlings
201 expressing GFP-ATG8e fusion protein (Xiong et al., 2007) were treated with 50 μ M
202 ABA for 3 and 6 h and subjected to additional treatment of 200 μ M NaHS for 1 h. Total
203 protein extracts were obtained, and immunoblot analysis was performed using anti-GFP
204 antibodies to detect free GFP and the GFP-ATG8e fusion protein (Figure 2A). A clear
205 increase in the free GFP protein level in ABA-treated seedlings was detected compared

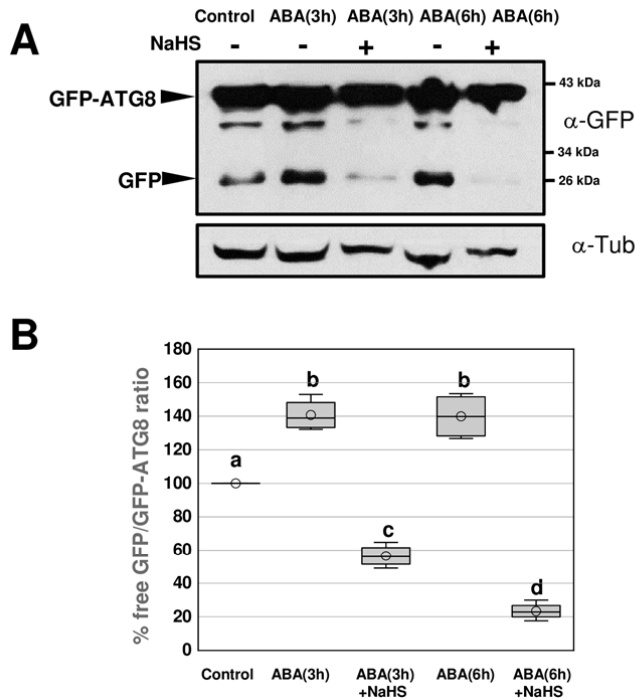


Figure 2. Effect of sulfide on autophagy induced by ABA treatment.

(A) Immunoblot analysis of GFP-ATG8e fusion protein. One-week-old seedlings expressing GFP-ATG8e fusion protein were transferred to liquid MS media and were not treated (control) or treated with 50 μ M ABA for 3 and 6 h and then 200 μ M NaHS for 1 h. Total protein extracts were prepared and subjected to SDS-PAGE and immunoblot analysis with anti-GFP antibodies. Anti-tubulin antibodies were used as the protein-loading control.

(B) Quantification of the free GFP/GFP-ATG8 ratio. For each condition, the levels of free GFP and GFP-ATG8e fusion protein were quantified in a less exposed blot shown in Supplemental Figure 6. The value of 100% was assigned to the free GFP/GFP-ATG8 ratio of the control sample. Data are from three independent experiments and evaluated by two-factor ANOVA. Same letters indicate no significant differences. $P < 0.05$.

206 with the control and a significant reduction in the GFP accumulation after the additional
 207 sulfide treatment was observed resulting in protein levels lower than those detected in
 208 the control. Quantification of the ratio free GFP/GFP-ATG8 under each condition
 209 showed that ABA induced the autophagic flux and that this ABA-induced autophagy
 210 was repressed by NaHS (Figure 2B). These findings and the proteomic data suggest that
 211 sulfide is a negative regulator of bulk autophagy independently of the condition used to
 212 induce this catabolic process, including at least nutrient limitation (Alvarez et al., 2012;
 213 Laureano-Marin et al., 2016a) and ABA-dependent stress (this study). Furthermore,
 214 persulfidation appears to be the mechanism of action of sulfide and the AtATG4a
 215 protease may be one of the specific targets.

216

217 **Persulfidation of AtATG4a at the Cys170**

218

219 To demonstrate the persulfidation-based modification of Cys residues in AtATG4a,
220 recombinant protein was purified, subjected to the tag-switch procedure, and analyzed
221 by immunoblotting using anti-biotin antibodies (Aroca et al., 2017a). A biotin-labeled
222 protein band corresponding to AtATG4a was clearly detected by the antibody.
223 Moreover, when AtATG4a was pretreated with DTT to reduce the persulfide residues,
224 the biotin-labeled protein bands were not detected (Figure 3A). These results clearly
225 indicate that AtATG4a undergoes persulfidation *in vitro*. To identify the Cys residue
226 that can be modified by persulfidation, recombinant AtATG4a was analyzed by liquid
227 chromatography (LC)-tandem mass spectrometry (MS/MS). The protein was digested
228 with trypsin under nonreducing conditions in the absence of alkylating agents to avoid
229 the reduction or modification of the persulfide residues. The digested peptides were
230 analyzed to detect a 32 Da molecular mass increase in the fragmentation spectrum. The
231 identified peptides included DTTYTSDVNWGCMIR that showed a persulfidation
232 modification of Cys170 (Figure 3B). AtATG4a protein was identified with a sequence
233 coverage of 97%, and no other persulfidated peptides were detected despite the presence
234 of other eleven Cys residues in the primary structure (Figure 3C). Cys170 is located in
235 the highly conserved catalytic site of ATG4 proteins from various organisms (Satoo et
236 al., 2009; Perez-Perez et al., 2014; Perez-Perez et al., 2016) suggesting that the
237 modification by persulfidation may have an important impact on the AtATG4a
238 proteolytic activity and biological function.

239

240 ***In Vitro* Processing of CrATG8 by AtATG4a**

241

242 To determine whether modification of AtATG4 by persulfidation has an effect on its
243 biological activity, an *in vitro* assay using CrATG8 from *C. reinhardtii* as a substrate
244 was established (Perez-Perez et al., 2010). CrATG8 was previously used to monitor the
245 activity of yeast ATG4 (Perez-Perez et al., 2010; Perez-Perez et al., 2014). ATG4
246 processes ATG8 at a conserved glycine residue located at the C-terminus of the protein
247 (Kirisako et al., 2000). Arabidopsis contains nine different ATG8 isoforms, none of
248 which has more than five amino acid residues after the conserved glycine (Doelling et
249 al., 2002). In contrast, CrATG8 harbors a 14-amino acid extension after the glycine, and
250 CrATG8 processing can be easily monitored by Coomassie Blue-stained SDS-PAGE

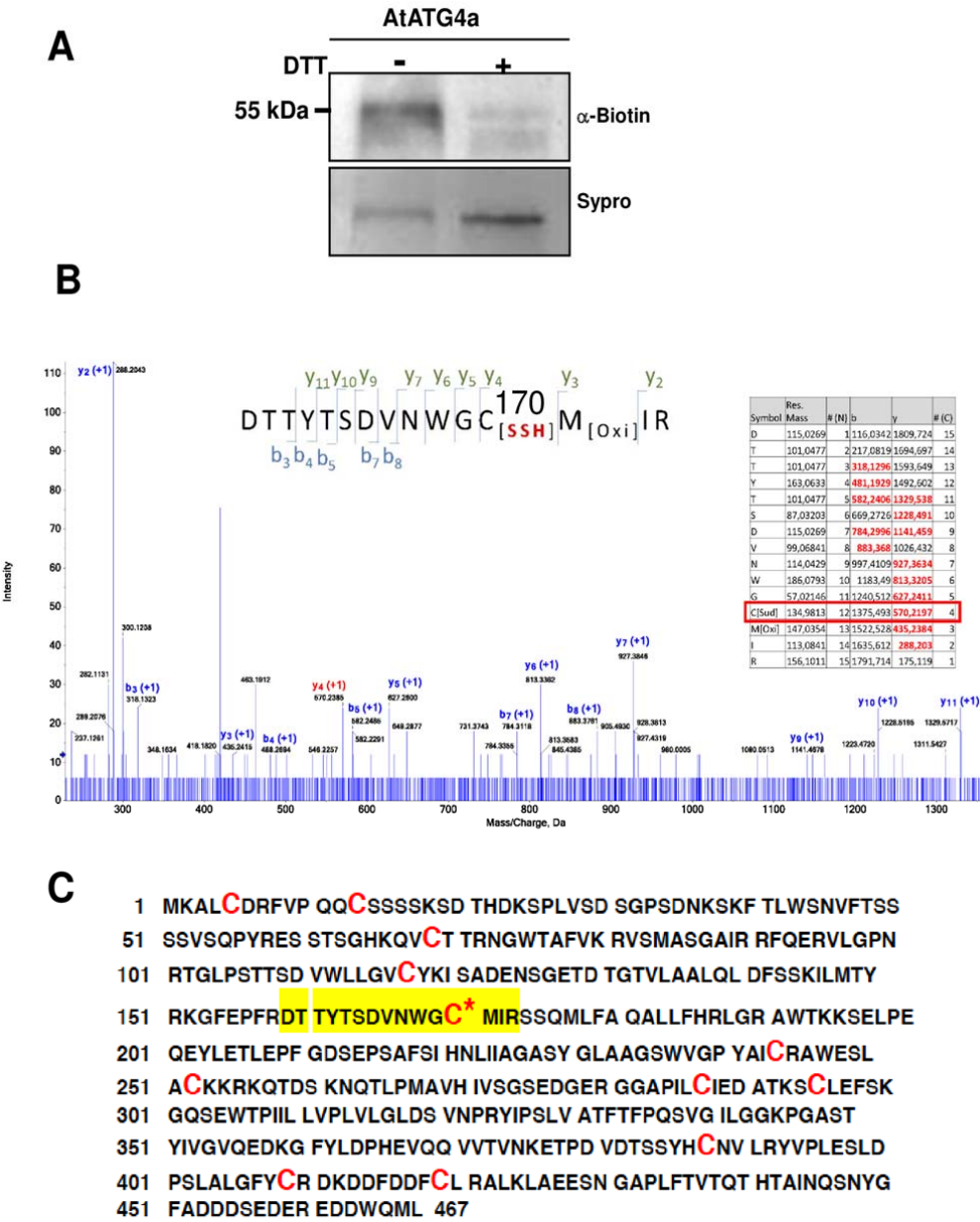


Figure 3. Persulfidation of Arabidopsis AtATG4a

(A) Immunoblot analysis of persulfidated AtATG4a. Purified recombinant AtATG4a was treated in the absence (-) or in the presence of 50 mM DTT (+) for 30 min at 4°C, dialyzed, and subjected to the tag-switch labeling as described in the Methods. Then, the proteins were subjected to immunoblot analysis using anti-biotin antibodies. Sypro Ruby fluorescent staining is shown as the protein loading control.

(B) Analysis of AtATG4a using mass spectrometry. LC-MS/MS analysis of a tryptic peptide of AtATG4a containing Cys170. The table inside the spectrum contains the predicted ion types for the modified peptide, and the ions detected in the spectrum are highlighted in red.

(C) AtATG4a protein sequence identified with 97% coverage. The peptide containing persulfidated Cys170 is highlighted in yellow and all the Cys residues are red.

due to differences in mobility between the unprocessed and processed CrATG8 forms (Supplemental Figure 1). When purified AtATG4a was incubated with CrATG8 in the

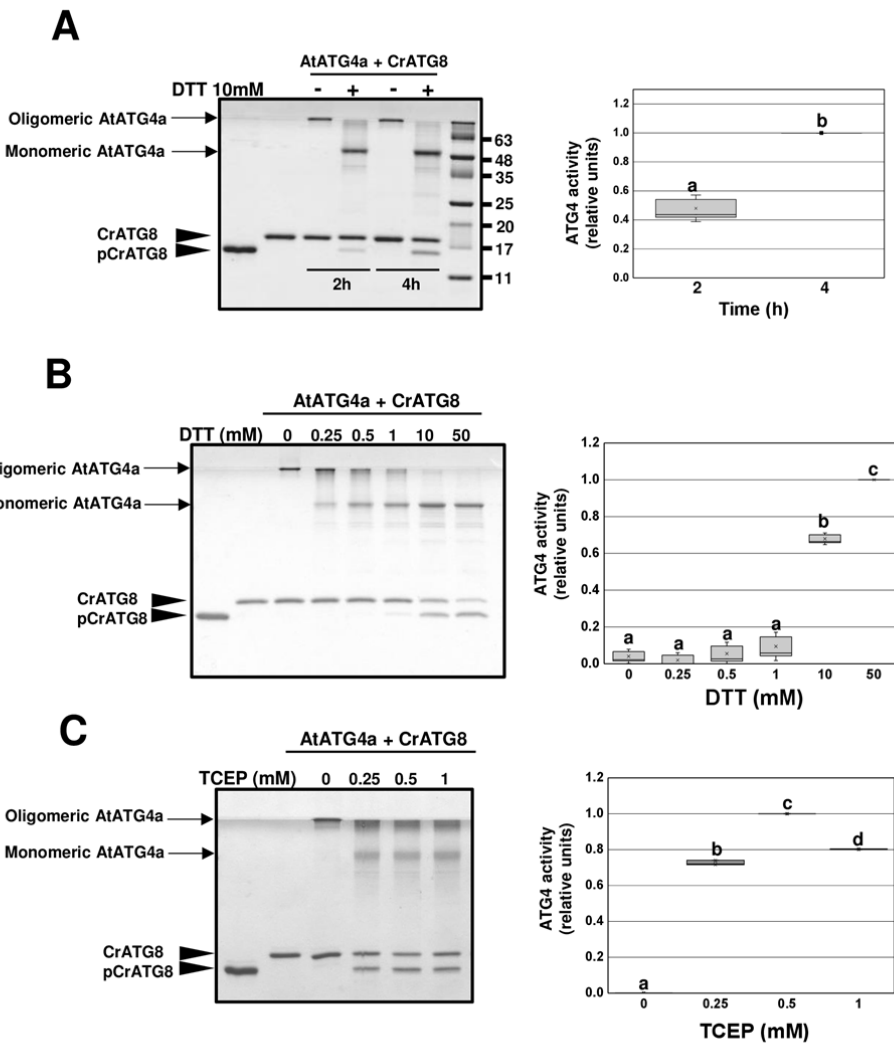


Figure 4. Arabidopsis ATG4a cleaves Chlamydomonas ATG8.

ATG4 activity of the recombinant AtATG4a was assayed by monitoring the cleavage of CrATG8 from the unprocessed (CrATG8) to processed (pCrATG8) forms (indicated by arrowheads) using SDS-PAGE followed by Coomassie Blue staining and quantification of protein band intensities. The ATG4 activity (relative units) was determined as the ratio of the band intensity of the processed CrATG8 to the sum of the intensities of the unprocessed and processed CrATG8. Activity value of 1 corresponds to the maximum. Processed pCrATG8 (lane 1) and unprocessed CrATG8 (lane 2) were loaded as controls. Representative images are shown. Data are from three independent experiments and evaluated by two-factor ANOVA. Same letters indicate no significant differences. P < 0.05.

(A) Effect of incubation time. AtATG4a was incubated with CrATG8 in the absence or in the presence of 10 mM DTT for the indicated times.

(B) Effect of DTT concentration. AtATG4a was incubated with CrATG8 in the absence or in the presence of increasing concentrations of DTT for 4 h.

(C) Effect of TCEP concentration. AtATG4a was incubated with CrATG8 in the absence or in the presence of increasing concentrations of TCEP for 4 h.

253 presence of DTT, the processed protein with the same mobility as a truncated form
 254 lacking the last 14 amino acids (CrATG8^{G120}, referred to as pCrATG8) was detected
 255 (Figure 4A). Therefore, AtATG4a was active and was able to process CrATG8 at its C-

256 terminal Gly validating the ATG4 processing activity assay. The results indicated that
257 AtATG4a activity was increased in a time-dependent manner and required a reducing
258 agent to adopt the monomeric form required for the activity (Figure 4A), as described
259 previously in other systems (Perez-Perez et al., 2010). To analyze the effect of a
260 reducing agent on AtATG4a activity, the *in vitro* assay in the presence or in the absence
261 of different DTT or TCEP concentrations was performed. In the absence of DTT and
262 TCEP, recombinant AtATG4a was in an oligomeric form that was retained in the upper
263 part of the acrylamide gel. Increasing concentrations of the reducing agent enhanced the
264 monomerization of recombinant AtATG4a and consequently the processing of CrATG8
265 by AtATG4a (Figure 4B and 4C). Therefore, properties of Arabidopsis ATG4a were
266 similar to those of the Chlamydomonas and yeast ATG4 (Perez-Perez et al., 2014;
267 Perez-Perez et al., 2016). Interestingly, TCEP was more efficient than DTT in the
268 activation of CrATG8 cleavage, showing the maximum level of ATG4 activity at 0.5
269 mM TCEP while two orders of magnitude higher DTT concentration was required for
270 optimal activity (Figure 4B and 4C).

271

272 **Sulfide inhibits the proteolytic activity of AtATG4a**

273

274 A suitable ATG4 enzyme activity assay was developed and used to study whether
275 persulfidation plays a role in the regulation of this activity. Thus, purified recombinant
276 AtATG4a was pretreated with TCEP to produce an active enzyme and then incubated
277 with increasing concentrations of NaHS as a sulfide donor. The ATG4 activity-
278 dependent CrATG8 processing was determined using the Coomassie-stained gel
279 method (Figure 5A). An increase in the accumulation of the unprocessed CrATG8 form
280 was detected when AtATG4a was incubated in the presence of NaHS at a concentration
281 as low as 0.1 mM. ATG4 activity was determined as the ratio of the band intensity of
282 the processed CrATG8 to the sum of the intensities of the unprocessed and processed
283 CrATG8; the data indicate that 0.1 mM NaHS significantly inhibits the ATG4 activity
284 to approximately 40% of the activity in the absence of sulfide (Figure 5B). Interestingly,
285 analysis of the aggregation state of AtATG4a in the presence of the sulfide donor
286 demonstrated that NaHS has different effects on monomerization and activity of
287 AtATG4a (Figure 5C). Sulfide did not promote the oligomerization of the active
288 monomeric AtATG4a to the same extent as it inhibited ATG4 proteolytic activity under
289 the conditions used in the assays. Concentrations of NaHS from 0.1 to 0.5 mM did not

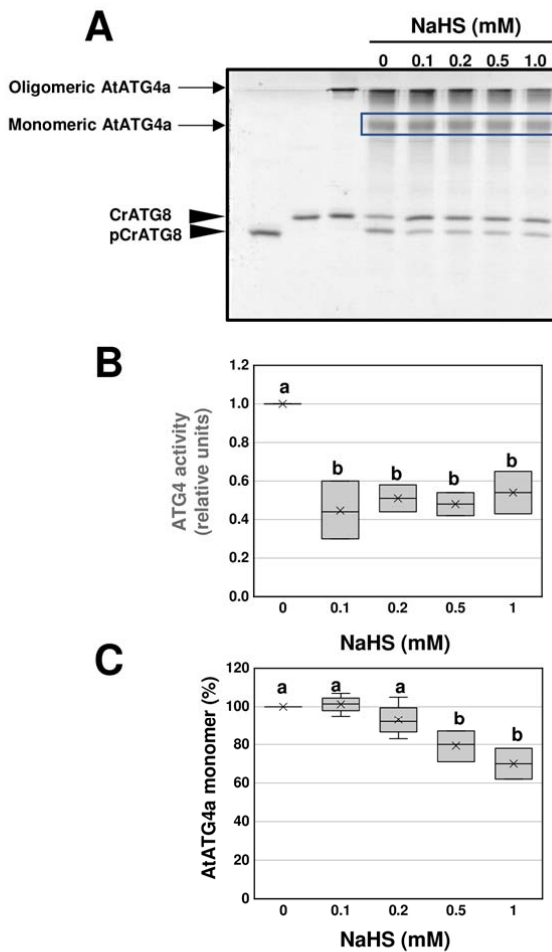


Figure 5. Effect of NaHS on AtATG4a enzyme activity.

(A) AtATG4a was incubated with 0.5 mM TCEP for 2 h and treated in the absence or the presence of indicated concentrations of NaHS for 1 h. Then, CrATG8 was added to the incubation mixture, and ATG4 activity was monitored after 4 h using Coomassie-stained gels as described in the Methods. All procedures were performed at 25°C. Lane 1 and 2, unprocessed and processed CrATG8, respectively. Lane 3, AtATG4a incubated with CrATG8 in the absence of TCEP and of NaHS. Lanes 4-8, TCEP-pretreated reduced AtATG4a incubated with CrATG8 in the presence of increasing concentrations of NaHS (from 0 to 1 mM). A representative image is shown.

(B) Quantification of ATG4 activity (relative units) determined as the ratio of the band intensity of the processed CrATG8 to the sum of the intensities of the unprocessed and processed CrATG8. A value of 1 corresponds to AtATG4a in the absence of NaHS (lane 4).

(C) Quantification of the protein band intensity corresponding to the monomeric AtATG4a form marked by a rectangle in A. A value of 100% corresponds to AtATG4a in the absence of NaHS (lane 4).

Data are from three independent experiments and evaluated by two-factor ANOVA. Same letters indicate no significant differences. P < 0.05

290 significantly increase the abundance of high molecular weight and inactive oligomers of
 291 AtATG4a (Figure 5C). These results indicate that sulfide donor inhibits AtATG4a
 292 activity but does not directly influence the aggregation state of the protein in contrast to
 293 the effect of DTT or thioredoxin on yeast and *C. reinhardtii* ATG4 proteins (Perez-
 294 Perez et al., 2014; Perez-Perez et al., 2016).

Recent studies have questioned whether NaHS can be the sulfurating molecule instead of the proposed polysulfide and persulfide molecules. These molecules contain sulfane sulfur, the form of sulfur (S^0) with the ability to reversibly attach to other sulfur atoms. Most of the reported biological activity associated with sulfide may be due to persulfides, which are considered responsible for the intracellular signal transduction through persulfidation in the living body (Toohey, 2011; Ida et al., 2014; Mishanina et al., 2015). Thus, assays similar to those described above were performed using various concentrations of polysulfide Na_2S_4 used as a sulfur donor. Our results indicated that polysulfide inactivates AtATG4a more efficiently than NaHS (Figure 6A and 6B). Concentrations of Na_2S_4 three orders of magnitude (10-50 μM) lower than those of NaHS were necessary to achieve a similar inhibition, and complete inactivation of the enzyme was observed at 100 μM Na_2S_4 . Furthermore, polysulfides were more active in promoting the aggregation state of AtATG4a, and the differences in the effects on activity and oligomerization were not as pronounced as those observed with NaHS (Figure 6C).

Collectively, our results indicate that sulfide can inhibit the ATG4 proteolytic activity and that this inhibition is independent on the ATG4 aggregation state, at least at low concentrations of sulfurating species. Our results also suggest that this inhibitory effect is mediated by specific persulfidation of the catalytic Cys170 residue. Site-directed mutagenesis of this catalytic cysteine inactivates ATG4 activity in all tested systems and conditions (Scherz-Shouval et al., 2007; Shu et al., 2010; Perez-Perez et al., 2014; Perez-Perez et al., 2016), and therefore the mutant enzyme cannot be used to test the inhibitory effect of sulfide.

To examine the impact of posttranslational modification of Cys170 by persulfidation on the interaction between AtATG4a and its substrate AtATG8a, 3D homology modelling was performed using the structure of the *Homo sapiens* HsATG4b-LC3 complex (Satoo et al., 2009). AtATG4a shares up to 33.4% sequence identity with HsAtg4B (E-value 7.3×10^{-90}) and AtATG8a shares up to 59.4% sequence identity with HsLC3 (E-value 1.8×10^{-40}) with conserved residues covering the whole sequence. Persulfidation of Cys170 in the AtATG4a-AtATG8a protein complex caused conformational changes and intramolecular rearrangements of the catalytic site (Figure 7A) influencing substrate recognition. Addition of the -SH group to Cys170 induced an unfavorable steric effect particularly affecting the His366 residue since the imidazole

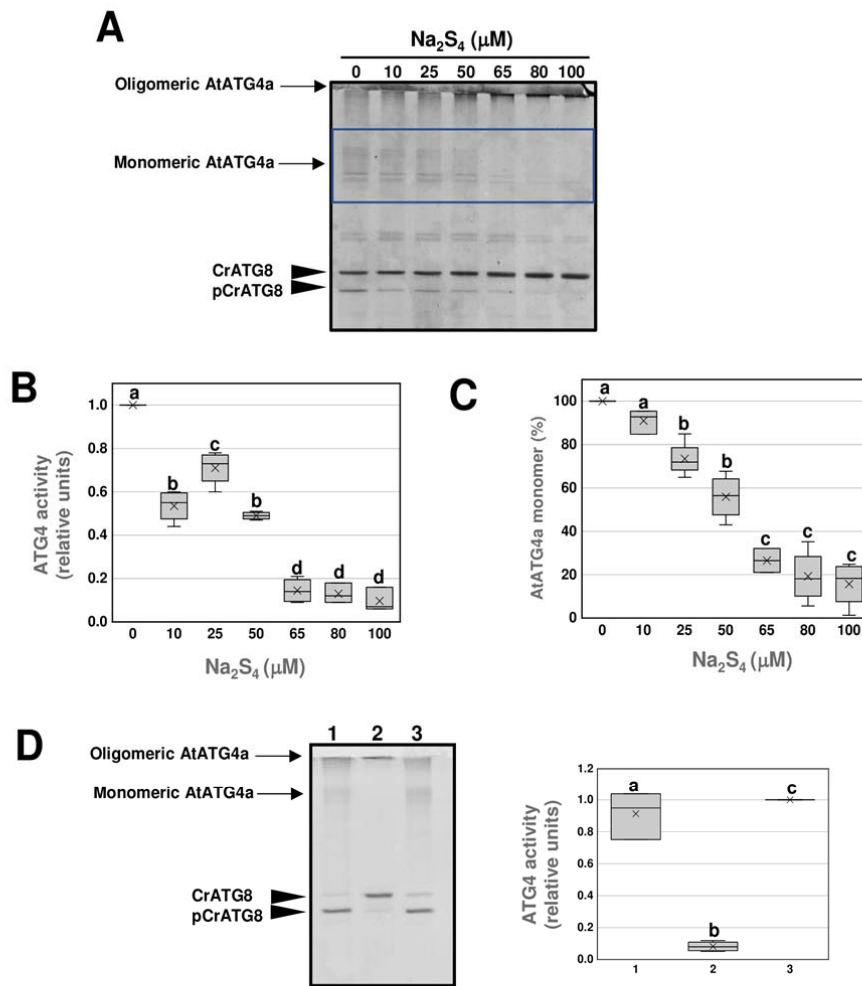


Figure 6. Effect of polysulfides on AtATG4a enzyme activity.

(A) AtATG4a was incubated with 0.5 mM TCEP for 2 h and subsequently treated in the absence or in the presence of indicated concentrations of Na₂S₄ for 1 h. Then, CrATG8 was added to the incubation mixture, and ATG4 activity was monitored after 4 h using Coomassie-stained gels as described in Methods. All procedures were performed at 25°C. A representative image is shown.

(B) Quantification of ATG4 activity (relative units) determined as the ratio of the band intensity of the processed CrATG8 to the sum of the intensities of the unprocessed and processed CrATG8. A value of 1 corresponds to AtATG4a treated in the absence of Na₂S₄ (lane 1).

(C) Quantification of the protein band intensity corresponding to the monomeric AtATG4a form marked by a rectangle in A. A value of 100% corresponds to AtATG4a treated in the absence of Na₂S₄ (lane 1).

(D) Reversibility of the effect. AtATG4a was first incubated with 0.25 mM TCEP for 2 h (lane 1) and treated with 100 mM Na₂S₄ for 1 h (lane 2) and 1 mM TCEP for 1 h (lane 3). ATG4 activity was monitored using Coomassie-stained gels. The experiment was performed at least three times and a representative image used for the quantification of the activity is shown.

Data are from three independent experiments and evaluated by two-factor ANOVA. Same letters indicate no significant differences. P < 0.05

328 Cδ2 and Nε2 atoms are 2.9 Å and 3.4 Å from the S atom of Cys170, respectively. The
 329 additional sulfur and hydrogen atoms have a covalent radius of 1.02 Å and 0.37 Å,
 330 respectively (Figure 7B). Therefore, this cysteine modification can inhibit the activity of

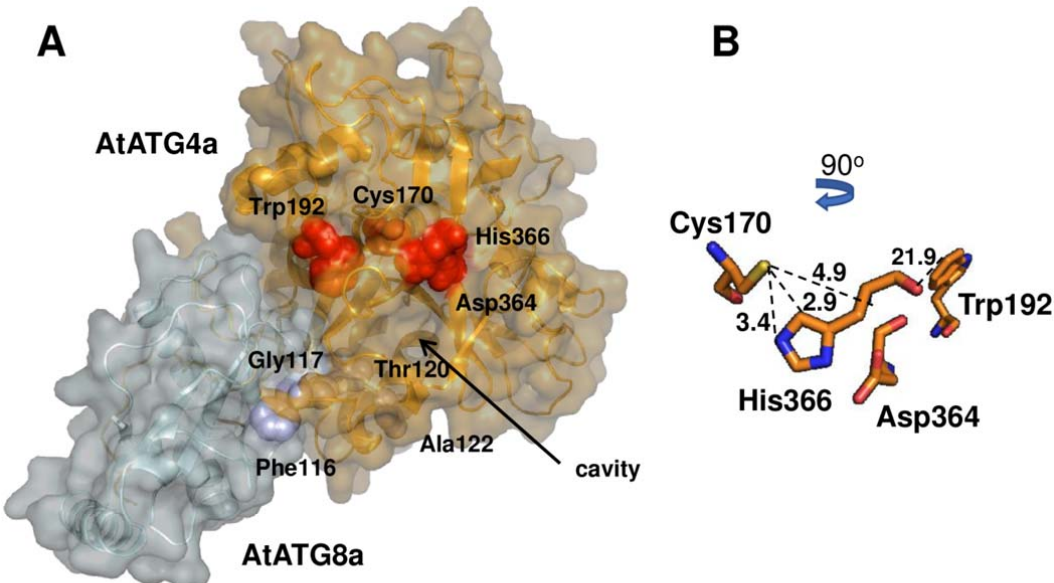


Figure 7. Predicted structure of the AtATG4a-AtATG8a complex.

(A) 3D modelling of the AtATG4a-AtATG8a complex based on the structure of the HsAtg4B-LC3 protein complex (PDB ID: 2Z0E). The AtATG8 protein sequence (Q8LEM4 in UniProtKB) corresponds to the splice variant 1. Surface representation of the protein complex and the equivalent residues surrounding the catalytic cavity Cys170, Trp192, Asp364, and His366 in AtATG4a (red) and Phe116, Gly117, Thr120, and Ala122 in AtATG8a (blue) are shown in the structural models.

(B) Zoomed view of the putative conformation of the active site showing the spheres corresponding to the position and distance (\AA) of catalytic residues Cys170, Trp192, Asp364, and His366 in AtATG4a.

331 the plant AtATG4a protein. Interestingly, the simulation of surface electrostatic
 332 potential reveals the differences between HsATG4b-LC3 and AtATG4a-AtATG8a
 333 protein complexes; specifically, the electrostatic potential surface around the catalytic
 334 site of the human complex is more electronegative compared to that of the Arabidopsis
 335 complex (Supplemental Figure 2). However, cysteine persulfidation does not perturb
 336 the charge, since the additional $-\text{SH}$ group is a neutral contribution. A closer look at the
 337 catalytic cavity region suggests that it is somewhat smaller and less exposed to the
 338 solvent in Arabidopsis compared to the human homologue likely hampering the
 339 accessibility of the substrate. Overall, these data suggest that inhibition of the
 340 processing activity of Arabidopsis AtATG4a may be due to a conformational change of
 341 the catalytic site induced by persulfidation.

342

343 **Endogenous Arabidopsis ATG4 processes CrATG8 for conjugation**

344

345 Our results indicate that sulfide can regulate ATG4 proteolytic activity through
346 persulfidation of a specific cysteine residue. This conclusion is based on *in vitro*
347 experiments. If sulfide has a biological role in the control of ATG4 activity in living
348 plants, the inhibitory effect of sulfide should be reversible. To explore this hypothesis,
349 active AtATG4a was inhibited by a high concentration of Na₂S₄ to promote complete
350 inactivation and to determine the extent of reversion by reduction of all persulfide
351 residues with TCEP. The reactivation of AtATG4a was monitored as the ratio of the
352 protein band intensities of unprocessed and processed CrATG8 (Figure 6D). The results
353 clearly confirm that the inhibition of ATG4 proteolytic activity by sulfide is reversible,
354 suggesting that it may play a role in the control of ATG4 *in vivo*.

355 To determine whether sulfide regulation of ATG4 also occurs *in vivo*, ATG4 enzyme
356 activity was detected in Arabidopsis leaves using the addition of exogenous CrATG8 to
357 leaf protein extracts. Immunoblot analysis using antibodies against CrATG8 indicated
358 that CrATG8 was efficiently processed by incubation with Arabidopsis leaf protein
359 extracts (Figure 8A, right part). To confirm the specificity of the CrATG8 cleavage
360 assay, Arabidopsis leaf protein extracts were incubated with a mutant of CrATG8 with
361 conserved Gly-120 replaced by Ala (G120A). This CrATG8 mutant is not processed by
362 ATG4 (Perez-Perez et al., 2010; Perez-Perez et al., 2016). When G120A was used as
363 the substrate, the processed form was not detected, demonstrating the specificity of the
364 endogenous ATG4 activity in Arabidopsis (Figure 8A, left part). Moreover, in addition
365 to full-length and processed CrATG8, the antibodies specifically recognized other bands
366 with faster mobility than pCrATG8 (Figure 8, asterisks). These bands were exclusively
367 detected when wild type CrATG8, but not the G120A mutant, was used in the assay and
368 they apparently correspond to the conjugated form of CrATG8 protein, as demonstrated
369 previously in Chlamydomonas (Perez-Perez et al., 2010). Incubation of the Arabidopsis
370 protein extract with the processed form of the CrATG8 (pCrATG8), which does not
371 require cleavage by ATG4 and can be conjugated to PE (Supplemental Figure 3),
372 confirmed that the faster mobility bands correspond to lipidated forms. The antibodies
373 detected mainly an intense protein band that was progressively accumulated with
374 incubation time and fully lipidated at the shortest time of incubation with the extracts
375 under autophagy-activating conditions induced by nitrogen deficiency (Supplemental
376 Figure 3). Therefore, our data demonstrated that the Arabidopsis protein extracts
377 contain all the active proteins required for the conjugation of ATG8 and efficiently

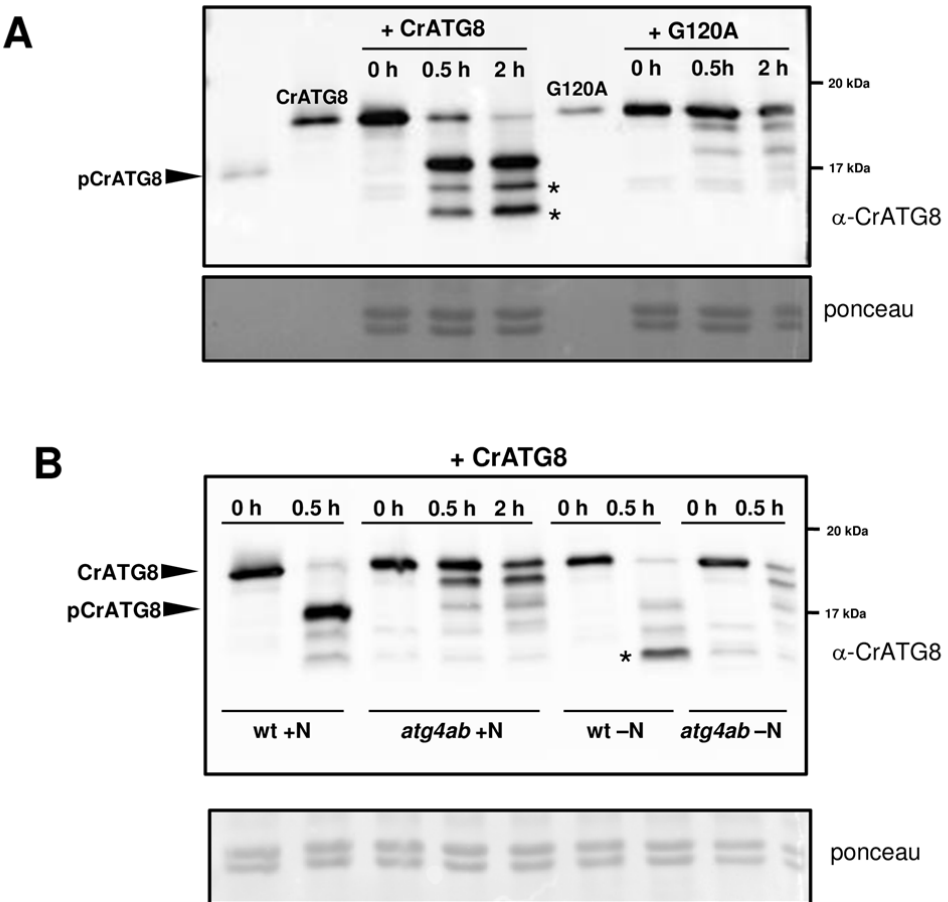


Figure 8. Cleavage and conjugation of Chlamydomonas ATG8 by Arabidopsis proteins.

(A) ATG4 proteolytic activity in wild-type Arabidopsis leaves. Arabidopsis protein extracts prepared from leaves of wild-type seedlings grown for 11 days on MS medium were incubated with CrATG8 or site-directed mutant G120A proteins at 25°C for the indicated times, and ATG4 activity was monitored as the cleavage of the ATG8 forms to the processed (pCrATG8) forms by immunoblotting analysis with anti-CrATG8. Processed pCrATG8 (lane 1), unprocessed CrATG8 (lane 2), and site-directed mutant G120A (lane 6) were loaded as controls.

(B) ATG4 proteolytic activity in the Arabidopsis *atg4ab* mutant. Arabidopsis protein extracts were prepared from the leaves of wild-type and *atg4ab* double mutant seedlings grown for 7 days on the MS medium and transferred to the same medium (+N) or to a nitrogen-deficient medium (-N) for additional 4 days. The protein extracts were incubated with CrATG8 at 25°C, and ATG4 activity was monitored after 0, 0.5, or 2 h, as indicated by immunoblotting analysis with anti-CrATG8.

The arrowheads show the unprocessed CrATG8 and processed pCrATG8 protein bands, and the asterisks indicate faster-mobility protein bands. Ponceau staining is shown as the protein loading control of the Arabidopsis extract.

378 recognize CrATG8, thus validating the results of the ATG4 activity assay in the cell-
379 free total extract.

380 To confirm the conclusion that Arabidopsis ATG4 proteins catalyze the processing
381 of CrATG8 in the cell-free total extract assay, the ATG4 enzyme activity of the
382 Arabidopsis protein extracts prepared from the *atg4ab* double-mutant seedlings was

assayed (Supplemental Figure 4) (Yoshimoto et al., 2004; Chung et al., 2010). When CrATG8 was incubated with the Arabidopsis *atg4ab* protein extract, the processed form of CrATG8 was not detected even after extended incubation. Similarly, when protein extracts were prepared from nitrogen-limited seedlings, the immunoblot analysis revealed a prominent protein band corresponding to the lipidated form of CrATG8 only in the presence of protein extracts from wild-type plants but not from the *atg4ab* mutant (Figure 8B).

Therefore, our data show that endogenous Arabidopsis ATG4 proteins recognize the cleavage site of the Chlamydomonas ATG8 substrate, which is efficiently processed in the cell-free enzymatic assay.

393

394 **Sulfide reversibly inhibits the ATG4 activity in Arabidopsis seedlings**

395

The effect of a sulfurating species on endogenous Arabidopsis ATG4 proteins was investigated to confirm the results obtained in *in vitro* analysis. Leaf protein extracts were treated with polysulfide, and CrATG8-processing activity was compared with that in the untreated extract. A pronounced decrease in the ATG4 activity was observed when the Arabidopsis extract was pretreated with Na₂S₄ (Figure 9A). Additionally, treatment of the Arabidopsis leaf protein extract with the alkylating agent iodoacetamide inhibited ATG4 activity as expected because ATG4 is a Cys protease and its activity is dependent on the catalytic Cys; this effect was similar to the effect of polysulfide (Figure 9A, left panel). Thus, these findings strongly suggest that sulfide negatively regulates ATG4 activity *in vivo*, at least the cleavage activity of the C-terminal extension of ATG8. Reversibility of the inhibitory effect of polysulfide was also analyzed. When the polysulfide-incubated Arabidopsis extract was treated with TCEP as a reducing agent to reduce the persulfide-modified cysteine residues in the protein extract, a significant difference in the activity was detected compared with that in the polysulfide-treated extract (Figure 9A, right panel). When the extract was incubated with Na₂S₄, the processed CrATG8 form was barely detected; however, incubation with TCEP significantly increased the abundance of this band, suggesting that sulfide may inhibit Arabidopsis ATG4 activity in a reversible manner.

To characterize the inhibition of ATG4 proteolytic activity by sulfide, the effect of polysulfide on the ATG4 activity was assayed under basal and autophagy-inducing conditions. Two different established conditions of autophagy induction were tested:

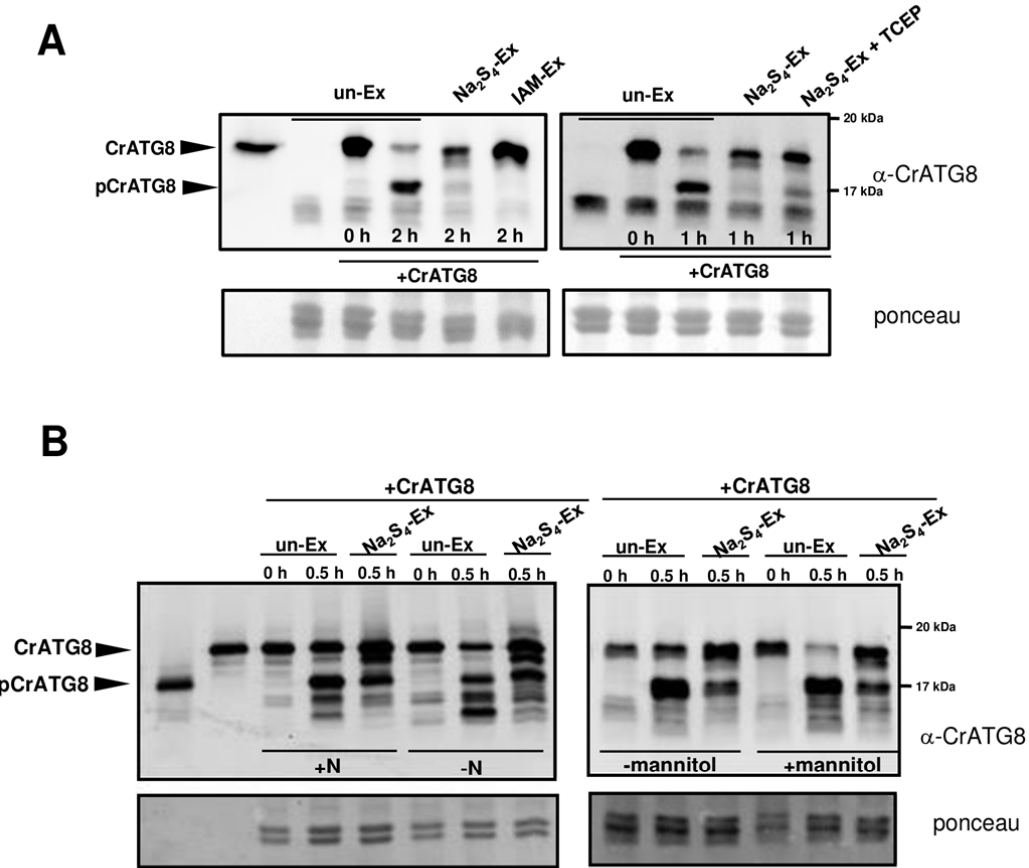


Figure 9. Sulfide inhibits the endogenous proteolytic activity of *Arabidopsis* ATG4.

(A) Effect of polysulfides on endogenous enzyme activity of *Arabidopsis* ATG4. *Arabidopsis* protein extracts (Ex) were prepared from the leaves of seedlings grown for 11 days on the MS medium. The extracts were treated in the absence (un-Ex) and in the presence of 200 μM Na_2S_4 ($\text{Na}_2\text{S}_4\text{-Ex}$), or 20 mM iodoacetamide (IAM-Ex) for 30 min, or in the presence of 200 μM Na_2S_4 for 30 min and 1 mM TCEP for 30 min ($\text{Na}_2\text{S}_4\text{-Ex} + \text{TCEP}$). Then, CrATG8 was added to the incubation mixture, and ATG4 proteolytic activity was monitored. Lane 1, unprocessed CrATG8.

(B) Sulfide reverts the endogenous enzyme activity of *Arabidopsis* ATG4 under autophagy-induced conditions. *Arabidopsis* protein extracts were prepared from the leaves of seedlings grown for 7 days on the MS medium and transferred to the same medium (+N) or to a nitrogen-deficient medium (-N) (left panel); or transferred to the same medium (-mannitol) or to same medium containing 300 mM mannitol (+mannitol) (right panel) for additional 4 days. The extracts were treated in the absence (un-Ex) or in the presence of 200 μM Na_2S_4 for 30 min ($\text{Na}_2\text{S}_4\text{-EX}$); CrATG8 was added to the incubation mixture and ATG4 activity monitored. Lane 1, processed pCrATG8 and lane 2, unprocessed CrATG8.

The ATG4 activity was monitored at the indicated times by immunoblotting analysis with anti-CrATG8. All procedures were performed at 25°C. Ponceau staining is shown as the protein loading control.

417 nitrogen deficiency, which has been extensively characterized previously (Doelling et
418 al., 2002; Hanaoka et al., 2002; Xiong et al., 2005; Phillips et al., 2008; Guiboileau et
419 al., 2013; Laureano-Marin et al., 2016aa), and osmotic stress (Liu et al., 2009) imposed

420 by the addition of mannitol to the growth medium, which induces ABA-dependent
421 signaling pathway. The processed form of CrATG8 was detected in total extracts under
422 the control and autophagy-activating conditions, and sulfide inhibited endogenous
423 ATG4 activity under both conditions (Figure 9B). The faster-mobility protein band
424 corresponding to the lipidated form of CrATG8 was more prominent under nitrogen
425 limitation than osmotic stress; however, sulfide treatment decreased the accumulation of
426 the conjugated forms of CrATG8 under both conditions.

427

428

430 **DISCUSSION**

431

432 Accumulating evidence emphasizes the importance of autophagy in plant growth and
433 development. This catabolic process is highly dynamic and occurs at the basal levels to
434 maintain cellular homeostasis during growth; however, autophagy is fine-tuned to
435 adjust plant metabolism to internal and external perturbations. Various regulators of
436 autophagy have been identified in plant systems, such as energy sensor Snf1-related
437 protein kinase 1 (SnRK1), target of rapamycin (TOR) kinase, the TOR downstream
438 substrate ATG1 kinase complex, and the ER stress sensor inositol-requiring enzyme-1
439 (IRE1) (Marshall and Vierstra, 2018b; Soto-Burgos et al., 2018).

440 Other regulators of autophagy, such as hydrogen sulfide, have been identified
441 although the details of the molecular mechanism of action of these regulators remain
442 poorly understood. In the animal systems, interactions of sulfide with autophagy have
443 been described in various pathologies; depending on the disease, sulfide can either
444 suppress or activate autophagy. In all cases, hydrogen sulfide has protective effects
445 (Sen, 2017; Wang et al., 2017; Wu et al., 2018). Despite substantial progress, the exact
446 mechanism of autophagy regulation by sulfide in mammals remains unknown. In plants,
447 particularly in *Arabidopsis*, the interplay between sulfide and autophagy has been
448 shown, and progress in understanding of the mechanism has been obtained. Hydrogen
449 sulfide generated in the cytosol functions as a signaling molecule negatively regulating
450 autophagy independently of the nutritional conditions. Furthermore, sulfide was shown
451 to repress autophagy via a mechanism that is independent of redox conditions (Alvarez
452 et al., 2012; Laureano-Marin et al., 2016a).

453 Sulfur and autophagy are also linked by the mechanism of *Arabidopsis* sensing of the
454 sulfur-containing amino acid cysteine. Two different pathways have been identified for
455 sensing of its carbon/nitrogen precursor and its sulfur precursor. The sulfur precursor is
456 transduced to TOR by downregulation of glucose metabolism; therefore, sulfide
457 increases glucose levels and TOR kinase activity, downregulating autophagy (Dong et
458 al., 2017). This study demonstrated that cytosolic sulfide is not the signal responsible
459 for the regulation but does not exclude chloroplast sulfide as the signaling molecule. In
460 fact, a proteomic study showed that other sulfurating species in addition to the cytosolic
461 sulfide can be responsible for regulation of diverse biological processes (Aroca et al.,
462 2017a).

463 The data of the present study emphasize sulfide regulation of autophagy. Our
464 findings indicate that ABA activates autophagy and hydrogen sulfide downregulates
465 this catabolic process. A link between autophagy and ABA was previously described
466 through the *Arabidopsis* multistress regulator TSPO, a tryptophan-rich sensory protein-
467 related. ABA induces TSPO as a heme scavenger, which binds the excessive or
468 deleterious heme and then is targeted for degradation through autophagy (Vanhee and
469 Batoko, 2011; Vanhee et al., 2011). Other connections between autophagy and ABA
470 signaling via TOR have been described. Under nonstressed conditions, TOR
471 phosphorylation of ABA receptors leads to inactivation of SnRK2 kinases and disrupts
472 ABA signaling. Under abiotic stress conditions, ABA activates SnRK2 that
473 phosphorylates RAPTOR resulting in the inhibition of TOR and consequential
474 activation of autophagy (Wang et al., 2018).

475 Previous studies have shown that the main mechanism of action of sulfide involves
476 persulfidation of reactive cysteine residues of the target proteins resulting in changes in
477 enzyme structure, activity and subcellular localization previously demonstrated in
478 several target plant proteins (Aroca et al., 2015; Aroca et al., 2017b; Aroca et al., 2018).
479 A proteomic analysis performed on mature leaves from *Arabidopsis* plants grown under
480 nonstress conditions revealed that a high proportion of the whole *Arabidopsis* proteome
481 may undergo persulfidation under the basal conditions (Aroca et al., 2017a). In this
482 study, a comparative and quantitative proteomic analysis has been performed on ABA-
483 treated leaves to determine whether persulfidation mechanism is involved in sulfide
484 regulation of the ABA signaling pathways. Significant differences in the levels of
485 persulfidation were observed after ABA treatment compared to that under the control
486 conditions. Surprisingly, the protein with the lowest level of persulfidation after 3 h
487 ABA treatment was identified as Cys-type protease AtATG4a (5.20-fold change,
488 p<0.05, control *versus* 3h ABA), and even after 6 h of ABA treatment, the reduction in
489 the level of persulfidation remained very significant (2.46-fold change, p<0.05, control
490 *versus* 6 h ABA). These data suggest that AtATG4a may be a target of persulfidation
491 and that this posttranslational modification may be the molecular mechanism mediating
492 sulfide regulation of autophagy. A detailed analysis of the ABA-triggered changes in
493 the persulfidation proteome will be performed in a future study. Our results
494 demonstrated that ATG4 proteolytic activity in *Arabidopsis* is reversibly regulated by
495 sulfide, and this regulation effectively controls the progression of autophagy. Thus, our

496 findings contribute to the understanding of the mechanism of regulation of autophagy
497 by sulfide in the plant systems and add another level of regulation to this process.

498 The AtATG4a protease contains a specific site of persulfidation detected by the tag-
499 switch procedure and confirmed by mass spectrometry. The specifically persulfidated
500 cysteine residue is Cys170, which is a part of the characteristic catalytic triad Cys-His-
501 Asp of cysteine proteases (Sugawara et al., 2005). Comparison of the amino acid
502 sequences of ATG4s from various biological systems indicated low similarity although
503 the amino acids required for the cysteine protease activity, including the catalytic
504 Cys170, are highly conserved. Interestingly, Cys170 is the only Cys residue that is
505 conserved in all known ATG4s (Supplemental Figure 5) (Perez-Perez et al., 2014;
506 Perez-Perez et al., 2016; Seo et al., 2016). In contrast, redox regulation of specific Cys
507 residues has been detected in ATG4 from human, yeast, and Chlamydomonas.
508 However, these residues are not conserved in various organisms and therefore the
509 details of the regulatory mechanisms may be different. In humans, HsATG4a and
510 HsATG4b proteases are the targets of reversible oxidation by H₂O₂, and the cysteine
511 residue Cys81 located close to the catalytic Cys residue (Cys77, analogous to Cys170 in
512 Arabidopsis) was shown to be critical for this regulation (Scherz-Shouval et al., 2007).
513 Recently, Cys292 and Cys361 have been shown to be HsATG4b sites essential for
514 reversible oxidative modification (Zheng et al., 2020). Redox regulation of the yeast
515 ScATG4 protein is due to the formation of a disulfide bond between the noncatalytic
516 residues Cys338 and Cys394, which is thioredoxin-dependent (Perez-Perez et al., 2014).
517 In *Chlamydomonas reinhardtii*, CrATG4 activity depends on the formation of a single
518 disulfide bond regulated by the NADPH/thioredoxin system; however, only Cys400,
519 which is the equivalent to Cys338 in yeast, has been demonstrated to be required for
520 redox regulation of the algal CrATG4 enzyme (Perez-Perez et al., 2016). In
521 Arabidopsis, the activity of AtATG4a and AtATG4b is reversibly inhibited *in vitro* by
522 reactive oxygen species (Woo et al., 2014), although redox regulation of the critical
523 cysteine residues was not reported. Additionally, other posttranslational modifications
524 of cysteine residues of ATG4 proteases have been described, such as S-nitrosylation of
525 HsATG4b at Cys189 and Cys292, though these two residues are not conserved in the
526 HsATG4a amino acid sequence (Li et al., 2017). Therefore, persulfidation of the
527 catalytic Cys residue of cysteine proteases ATG4 can be a posttranslational
528 modification conserved in various biological systems.

529 Because the target residue of persulfidation involves the active site, an effect of the
530 sulfide donor molecules on the enzymatic activity of AtATG4a was anticipated. In fact,
531 a 3D modelling analysis predicted that the addition of a sulfur atom to the -SH group of
532 Cys170 can cause unfavorable effects on the catalytic site of AtATG4a that should
533 affect substrate recognition and impair the enzyme activity. To test this hypothesis, a
534 heterologous activity assay was developed using CrATG8 as a substrate similar to
535 previously reported assays (Perez-Perez et al., 2014; Seo et al., 2016). Our results
536 indicate that the plant protease is functional and can process the algal substrate.

537 Our findings demonstrate that sulfide plays a specific role in the regulation of ATG4
538 enzymatic activity. The sulfide donor molecules used in this study, such as hydrosulfide
539 and tetrasulfide, significantly inactivate AtATG4a cleavage activity even at relatively
540 low concentrations, being polysulfide the most efficient inhibitor. The chemical
541 mechanism of the reaction of the thiol groups with the sulfide molecule remains a
542 matter of debate. H₂S cannot directly react directly with thiols, and the cysteine group
543 must be partially oxidized (converted to disulfide, glutathiolated, S-nitrosylated, or to
544 sulfenic acid) prior to sulfide attack. Alternatively, certain chemical studies have
545 demonstrated that sulfane sulfur (S⁰) of the polysulfide molecule is responsible for the
546 production of persulfide during interaction with the thiol group (Toohey, 2011; Kimura,
547 2015; Mishanina et al., 2015). This phenomenon can explain why polysulfide is a more
548 potent inhibitor of ATG4 activity than hydrosulfide. Moreover, the inhibitory effect of
549 low concentrations of sulfide on AtATG4a activity compatible with the *in vivo*
550 conditions is reversible by a reducing agent suggesting a biological role of this effect.
551 Interestingly, an enzymatic mechanism of reversing persulfidation by the thioredoxin
552 system has been demonstrated in animal systems (Wedmann et al., 2016; Doka et al.,
553 2020). Thus, it is plausible that the thioredoxin system also functions in plants to
554 modulate the activity of AtATG4.

555 The biological significance of sulfide regulation of autophagy through reversible
556 inhibition of ATG4 protease is reinforced by the assays in *Arabidopsis* leaves under
557 basal and autophagy-inducing conditions. Our experimental system was shown to be
558 suitable for specific assay of *Arabidopsis* ATG4 processing activity by using various
559 experimental approaches. Endogenous plant ATG4 cysteine proteases specifically
560 recognize the Gly120 cleavage site in the substrate from green algae based on the
561 experiment with the G120A mutant of the CrATG8 substrate. The processed form was
562 not detected when the *Arabidopsis* protein extracts were deficient in ATG4a and

563 ATG4b enzymes, demonstrating the specificity of our experimental system. The
564 endogenous Arabidopsis ATG4 proteins mimic the effect of sulfurating molecules on
565 AtATG4a *in vitro*, including significant inhibition by the sulfide donor and reversal by a
566 reducing agent. A significant increase in the overall ATG4 protease activity in
567 Arabidopsis was detected under autophagy-inducing conditions, including nitrogen
568 starvation and osmotic stress. Thus, a correlation was detected between the progress of
569 autophagy and the ATG4 enzymatic activity estimated using the heterologous assay
570 method. Additionally, the inhibitory effect of sulfide on the protease activity was
571 observed under the conditions of induced autophagy. Overall, our findings suggest that
572 negative regulation of the progress of autophagy by sulfide is mediated by specific
573 persulfidation of cysteine protease ATG4. However, additional targets of sulfide cannot
574 be ruled out and further analysis is needed.

575 In conclusion, our data suggest a new level of regulation of ATG4 activity by sulfide.
576 Based on our findings, we propose that intracellular sulfide maintains high levels of
577 persulfidation of the ATG4 pool under basal conditions, resulting in the inhibition of
578 ATG4 proteolytic activity. ATG4 inhibition limits the formation of ATG8-PE adducts
579 and consequently *de novo* synthesis of autophagosomes. An increase in the intracellular
580 level of ABA transiently reduces the level of persulfidation of the ATG4 population and
581 then activates the protease activity of the enzyme to process ATG8 that can be further
582 lipidated (Figure 10).

583

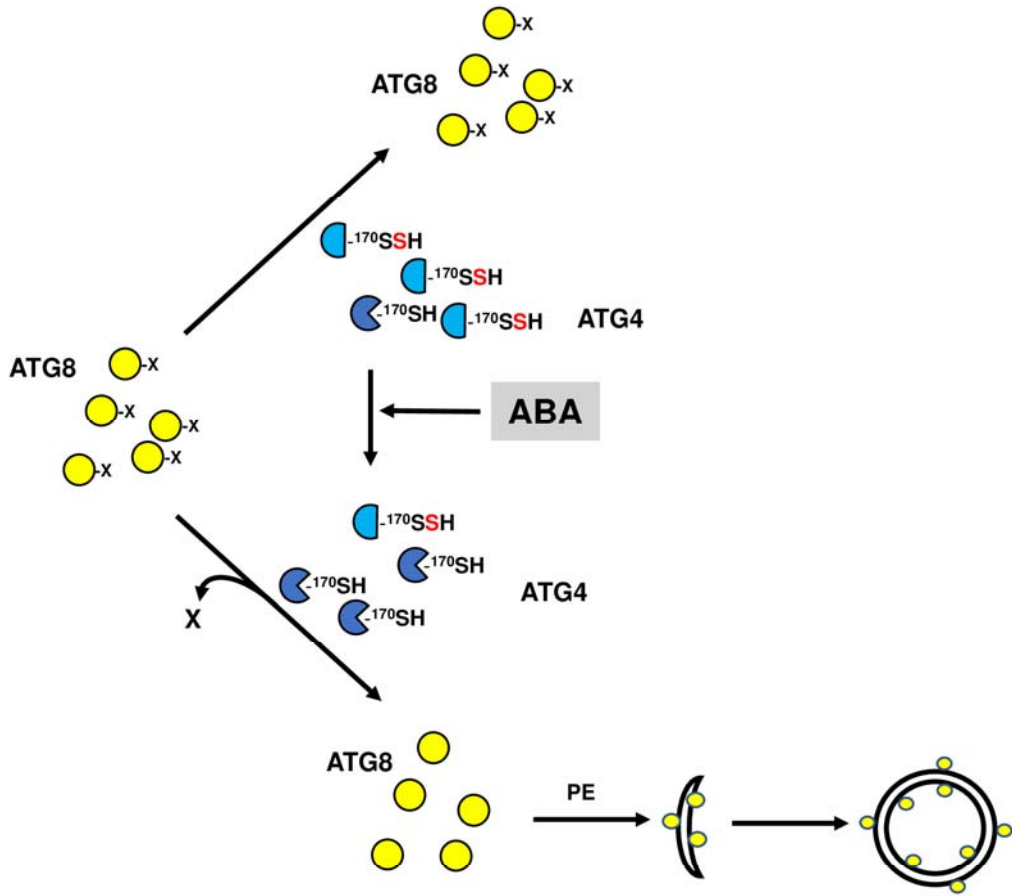


Figure 10. Graphical model of ABA-triggered autophagy induction through ATG4 posttranslational modification. Under basal conditions intracellular sulfide maintains high levels of persulfidation of the ATG4 pool which inhibits the proteolytic activity of the enzyme for ATG8 C-terminal processing. Increase in the intracellular level of ABA transiently decreases the level of persulfidation of the ATG4 population, activating the protease activity of the enzyme and the processing of ATG8 that can be further lipidated for the progression of autophagy. Yellow circles represent ATG8 protein with or without the C-terminal processed end (represented as X). Blue semicircles represent persulfidated ATG4 protein at the thiol ¹⁷⁰Cys residue. Blue Pacman symbols represent ATG4 protein with reduced thiol ¹⁷⁰Cys residue. The conjugation process of ATG8 with phosphatidylethanolamine (PE), autophagosome initiation and closure are also shown.

585 **METHODS**

586

587 **Plant Material, Treatments and Protein Extraction**

588

589 Plant growth conditions were 16 h of light ($120 \mu\text{E m}^{-2} \text{s}^{-1}$) at 22°C and 8 h of dark at
590 20°C . The *Arabidopsis thaliana* wild-type and the *atg4ab* mutant (Nottingham
591 Arabidopsis Stock Center) seeds were sown on the MS solid medium containing 0.8%
592 (w/v) agar, synchronized at 4°C for 2 d, and incubated vertically in a growth chamber.
593 For exposure to the nitrogen-starvation conditions, nitrogen-deficient MS medium was
594 prepared by replacing nitrate salts with chloride salts. For mannitol treatment, 300 mM
595 was added to the MS medium. One-week-old seedlings were transferred to nitrogen-
596 deficient or mannitol-containing MS solid medium for an additional 4 d of growth.

597 For proteomic analysis, wild-type *Arabidopsis* plants were grown in soil for 30 days,
598 and then sprayed with water (control conditions), or 50 μM ABA for 3 and 6 h. To
599 assess the autophagic activity, one-week-old *Arabidopsis* seedlings overexpressing
600 GFP-ATG8e (Xiong et al., 2007) were transferred to the MS liquid medium and treated
601 with 50 μM ABA for 3 and 6 h and with 200 μM NaHS for 1 h.

602 *Arabidopsis* material (200 mg) was ground in liquid nitrogen with 400 mL of
603 extraction buffer (100 mM Tris-HCl, pH 7.5, 400 mM sucrose, 1 mM EDTA, 0.1 mM
604 phenylmethylsulfonyl fluoride (PMSF)) using a mortar and pestle. After centrifugation
605 at 500 g for 10 min at 4°C , the supernatant was used as the *Arabidopsis* protein extract.

606

607 **Persulfidated Protein Quantitation by Label-free SWATH-MS Acquisition and**
608 **Analysis**

609

610 Protein samples from three biological replicates (independent pools) of leaf tissues
611 treated with ABA for 0 h (control sample), 3 h and 6 h were isolated and 1 mg of
612 protein per sample was used for the tag-switch labeling for enrichment of persulfidated
613 proteins as described (Aroca et al, 2017). After elution from the streptavidin-beads, the
614 proteins were precipitated by TCA/acetone. Precipitated samples were resuspended in
615 50 mM ammonium bicarbonate with 0.2% Rapigest (Waters) for protein determination.
616 Protein (50 μg) was digested with trypsin as described previously (García et al., 2019;
617 Vowinckel et al., 2014), and the SWATH-MS analysis was performed at the Proteomic

618 Facility of the Institute of Plant Biochemistry and Photosynthesis, Seville, Spain. A
619 data-dependent acquisition (DDA) approach using nano-LC-MS/MS was initially
620 performed to generate the SWATH-MS spectral library as described by García et al.,
621 (2019).

622 The peptide and protein identifications was performed using Protein Pilot software
623 (version 5.0.1, Sciex) with the Paragon algorithm. The search was conducted against a
624 Uniprot proteome (June 2017, release), and the corresponding reversed entries and
625 common contaminants were assembled in the FASTA format using ProteinPilot
626 software v5.0.1 (AB Sciex) with the Paragon™ algorithm. Samples were input as
627 unlabeled with no special factors, trypsin digested and MSBT alkylated. The
628 automatically generated report in ProteinPilot™ was manually inspected for FDR (false
629 discovery rate) cut-off proteins and only proteins identified at FDR \leq 1% were
630 considered for output and subsequent analysis.

631 For relative quantification using SWATH analysis, the same samples used to
632 generate the spectral library were analyzed using a data-independent acquisition (DIA)
633 method. Each sample (2 μ L) was analyzed using SWATH-MS acquisition method on
634 the LC-MS equipment with the LC gradient as described. The method consisted of
635 repeated acquisition cycles of TOF MS/MS scans (230 to 1500 m/z, 60 ms acquisition
636 time) of 60 overlapping sequential precursor isolation windows of variable width (1 m/z
637 overlap) covering the 400-1250 m/z mass range from a previous TOF MS scan (400-
638 1250 m/z, 50 ms acquisition time) for each cycle. The total cycle time was 3.7 s.

639 Autocalibration of the equipment and chromatographic conditions were controlled by
640 an injection of a standard of digested β -galactosidase from *E coli* between the replicates.

641 SWATH MS spectra alignment was performed with the *PeaKView* 2.2 (Sciex)
642 software with the *microapp SWATH* 2.0 using the reference spectral library generated as
643 described above. Two DIA raw files for each biological replicate were loaded in unison
644 using the following parameters: 10 peptides, 7 transitions, peptide confidence >99%,
645 1% FDR including shared peptides, and XIC width set at 0.05 Da. After data
646 processing, three distinct files were exported for subsequent quantification. The
647 processed mrkvw files containing protein information from PeakView were loaded into
648 MarkerView (Version 1.2.1, AB Sciex) for normalization of protein intensity (peak
649 area) for all runs using the built-in total ion intensity sum plug-in. Log2 transformation
650 was performed prior to statistical analysis. A histogram plot was used to check the
651 normality of distribution of each technical replicate. Mean values of protein expression

652 were used for calculation of fold change (FC). Proteins with adjusted $p < 0.05$ and
653 $FC \geq 1.5$ were considered differentially expressed.

654 The mass spectrometry proteomic data have been deposited to the ProteomeXchange
655 Consortium via the PRIDE (Vizcaino et al, 2016) partner repository with the identifier
656 PXD019802.

657

658 **Expression of AtATG4a in *Escherichia coli***

659

660 Total RNA was extracted from wild-type Arabidopsis leaves using an RNeasy plant
661 mini kit (Qiagen) and reverse-transcribed using an oligo (dT) primer and a SuperScript
662 first-strand synthesis system for RT-PCR (Invitrogen). Subsequently, a 1,404 bp
663 sequence encoding the full-length AtATG4a (At2g44140) protein was amplified by
664 PCR using the primers ATG4-F: CACCATGAAGGCTTATGTGA and ATG4-R:
665 ATGACTGGCAAATGCTCTGA and the proofreading Platinum Pfx DNA polymerase
666 (Invitrogen). The PCR conditions were as follows: a denaturation cycle of 2 min at 94°C
667 followed by 30 amplification cycles of 15 s at 94°C, 30 s at 57°C, and 1 min at 68°C.
668 The amplified cDNA was then ligated into the pENTR/D-TOPO vector using the
669 pENTR directional TOPO cloning kit (Invitrogen) according to the manufacturer's
670 instructions. Positive clones were identified by PCR and selected for plasmid DNA
671 isolation. The AtATG4a cDNA was then cloned into the expression vector pDEST17
672 using an *E. coli* expression system with gateway technology (Invitrogen), which
673 generates a fusion protein with an N-terminal 6x His tag that was confirmed by
674 sequencing; the expression was induced with L-arabinose in BL21-AI *E. coli* cells.

675

676 **Purification of the Recombinant AtATG4a Protein**

677

678 The 6x His-tagged recombinant protein was isolated from 200 mL of BL21-AI *E. coli*
679 cells that were cultured at 37°C to an optical density of 0.5 at 600 nm and induced with
680 0.2% (w/v) L-arabinose for 2.5 h at 37°C. Prior to purification, His-tagged AtATG4a
681 was solubilized with 6 M urea because the recombinant protein was contained in the
682 inclusion bodies. Then, the protein was purified from the soluble fraction by nickel
683 affinity chromatography using an Invitrogen Ni-NTA Purification System
684 (ThermoFisher Scientific), according to the manufacturer's instructions. The purified
685 protein was concentrated and desalting using 10 kD cutoff pore size centrifugal filter

686 units (Millipore). The purity of the protein was confirmed by SDS-PAGE using 12%
687 (w/v) polyacrylamide gels stained by Coomassie Blue.

688

689 **Detection of Persulfidation on the Recombinant AtATG4a**

690

691 An untreated aliquot of the purified recombinant AtATG4a and another aliquot
692 pretreated with 50 mM DTT for 30 min at 4°C to reduce all persulfide groups were
693 precipitated with acetone for 20 min at -20°C and centrifuged at a maximum speed for
694 20 min at 4°C. After acetone removal, the proteins were resuspended in 50 mM Tris-
695 HCl (pH 8) buffer supplemented with 2.5% (w/v) SDS and subjected to the tag-switch
696 procedure as described previously (Aroca et al., 2017a). The CN-biotinylated proteins
697 were then detected using an immunoblot assay as follows. The CN-biotinylated proteins
698 were separated using nonreducing SDS-PAGE through 12% (w/v) polyacrylamide gels
699 and transferred to polyvinylidene difluoride membranes (Bio-Rad) according to the
700 manufacturer's instructions. The anti-biotin (Abcam) and secondary antibodies were
701 diluted 1:500,000 and 1:100,000, respectively, and ECL Select western blotting
702 detection reaction (GE Healthcare) was used to detect the proteins using horseradish
703 peroxidase-conjugated anti-rabbit secondary antibodies. For protein loading control, the
704 membranes before immunodetection were stained with SYPRO Ruby (Invitrogen) to
705 detect all protein bands.

706

707 **Identification of Persulfidated Cys Residues of Recombinant AtATG4a Using
708 Mass Spectrometry**

709

710 Recombinant AtATG4a was separated using nonreducing SDS-PAGE through 12%
711 (w/v) polyacrylamide gels, and the band corresponding to AtATG4a was manually
712 excised from Coomassie-stained gels. Gel plugs were washed twice using 50 mM
713 ammonium bicarbonate and acetonitrile and dried under a steam of nitrogen. Then,
714 proteomics-grade trypsin (Sigma-Aldrich) at a final concentration of 16 ng/μl in 25%
715 (v/v) acetonitrile/50 mM ammonium bicarbonate solution was added; the samples were
716 digested at 37°C for 5 h. The reaction was stopped by adding 50% (v/v)
717 acetonitrile/0.5% (v/v) trifluoroacetic acid for peptide extraction. The eluted tryptic
718 peptides were dried by speed-vacuum centrifugation and resuspended in 6 μl of 0.1%

719 (v/v) formic acid in water. Digested peptides were subjected to nanoliquid
720 chromatography electrospray ionization tandem mass spectrometry analysis using a
721 nanoliquid chromatography system (ExcionLC AD, Sciex) coupled to a TripleTOF
722 5600+ mass spectrometer (Sciex) with a spray ionization source. Mass spectrometry and
723 MS/MS data of individual samples were processed using the Analyst TF 1.5.1 software
724 (Sciex). Peptide mass tolerance was set to $25 \mu\text{D D}^{-1}$ and 0.05 D for fragment masses,
725 and only one or two missed cleavages were allowed. Peptides with an individual M_r
726 search score ≥ 20 were considered correctly identified.

727

728 ***In vitro* ATG4 Enzyme Activity Assay**

729

730 The typical reaction mixture contained 5 μM recombinant AtATG4a, 5 μM CrATG8
731 (Perez-Perez et al., 2010), and 1 mM EDTA in Tris-buffered saline (50 mM Trizma
732 base, 138 mM NaCl, and 27 mM KCl at pH 8). When indicated, AtATG4a was
733 incubated in the presence of DTT, TCEP, NaHS, Na_2S_4 , or iodoacetamide alone or in
734 combination at the indicated times and concentrations. The reaction mixtures were
735 incubated at 25°C, and the reaction was stopped by the addition of β -mercaptoethanol-
736 free Laemmli sample buffer followed by 5 min boiling. The proteins were resolved
737 using nonreducing SDS-PAGE through 15% (w/v) polyacrylamide gels and stained
738 with Coomassie Brilliant Blue (Sigma-Aldrich). The gels were scanned with a GS-800
739 densitometer (Bio-Rad), and the signals corresponding to the unprocessed and
740 processed CrATG8 forms were quantified with the Quantity One software (Bio-Rad).
741 The ATG4 activity (in arbitrary units) was calculated as the ratio of the band intensity
742 of the processed CrATG8 to the sum of the intensities of the unprocessed and processed
743 CrATG8. An activity value of 1 corresponds to the maximum value.

744

745 **Assay of Endogenous ATG4 Enzyme Activity in Cell-Free Total Extract**

746

747 The *in vivo* assay of ATG4 activity in cell-free total extract was performed in a typical
748 reaction mixture containing 40 μg of leaf or 20 μg of root protein extract and 0.05 μM
749 purified unprocessed CrATG8, processed pCrATG8, or Gly-to-Ala mutant protein
750 (G120A). When required, a sulfur donor (Na_2S_4), a reducing agent (TCEP), or an
751 alkylating agent (IAM) was added at the indicated concentrations. The reaction mixture

752 was incubated at 25°C for the indicated time, stopped by addition of β-mercaptoethanol-
753 free Laemmli sample buffer, and boiled for 5 min. Then, proteins were separated using
754 nonreducing SDS-PAGE through 15% (w/v) polyacrylamide gels and transferred to
755 nitrocellulose membranes (Bio-Rad) as described previously (Perez-Perez et al., 2010).
756 Anti-CrATG8 (Pérez-Pérez et al., 2010) and secondary antibodies were diluted 1:3,000
757 and 1:10,000, respectively. An ECL Select western blotting detection reaction (GE
758 Healthcare) was used to detect the proteins. For protein loading control, the membranes
759 before immunodetection were stained with Ponceau S (Sigma-Aldrich) to detect all
760 protein bands.

761

762 **Protein modeling**

763

764 3D homology modeling was driven by Modeller (Sali and Blundell, 1993) using the
765 structure of the *Homo sapiens* Atg4B-LC3 complex (PDB ID: 2Z0E) (Sato et al., 2009)
766 as a template. Molecular crystal X-ray structures and structural model of *Arabidopsis*
767 *thaliana* AtATG4a-AtATG8a complex were inspected, analyzed and plotted with
768 PyMol 1.4.1 (Schrodinger LLC). Surface electrostatic potentials were calculated and
769 visualized using the PyMol 1.4.1 software.

770

771

772 **Accession Numbers**

773

774 The mass spectrometry proteomic data have been deposited to the ProteomeXchange
775 Consortium via the PRIDE partner repository with the identifier PXD019802. Sequence
776 data from this article can be found in the EMBL/GenBank data libraries under the
777 following accession numbers: *AtATG4a* (At2g44140), *AtATG4b* (At3g59950), *AtATG8a*
778 (At4g21980) and *CrATG8* (Cre16.g689650.t1.1).

779

780

781

782 **Supplemental Data**

783

784 **Supplemental Dataset 1.** Spectral library

785

786 **Supplemental Dataset 2.** Proteins quantified using SWATH acquisition method in the
787 control sample versus 3 h ABA treatment sample.
788

789 **Supplemental Dataset 3.** Proteins with different abundance in the control sample
790 compared to that in the 3 h ABA treatment sample at p value <0.05.
791

792 **Supplemental Dataset 4.** Proteins quantified using SWATH acquisition method in the
793 control sample versus 6 h ABA treatment sample.
794

795 **Supplemental Dataset 5.** Proteins with different abundance in the control sample
796 compared to that in the 6 h ABA treatment sample at p value <0.05.
797

798 **Supplemental Table 1.** Classification of the proteins with persulfidation level reduced
799 after 3 h ABA treatment.
800

801 **Supplemental Table 2.** Protein list of the elements included in the bin of "proteins"
802 with persulfidation level reduced after 3 h ABA treatment.
803

804 **Supplemental Figure 1.** C-Terminus of nine Arabidopsis and the Chlamydomonas
805 ATG8 proteins.
806

807 **Supplemental Figure 2.** Simulation of surface electrostatic potential distribution in the
808 HsATG4b-LC3 protein complex (PDB ID: 2Z0E) and AtATG4a-AtATG8a 3D
809 structural model.
810

811 **Supplemental Figure 3.** Conjugation of Chlamydomonas ATG8 processed by ATG4s
812 proteases from Arabidopsis.
813

814 **Supplemental Figure 4.** Phenotypes of wild-type and *atg4ab* double-mutant seedlings
815 under the basal and induced autophagy conditions.
816

817 **Supplemental Figure 5.** Alignment of amino acid sequences of ATG4 from various
818 sources.
819

820 **Supplemental Figure 6.** Lower exposition of the GFP-blot shown in Figure 2.

821

822

823

824

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826

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841 **AUTHOR CONTRIBUTIONS**

842

843 Á.A., A.M.L-M., M.E.P-P., I.Y., A. J-F. and I.M. performed research and analyzed the
844 data. J.L.C. and L.C.R. designed research and analyze data. C.G. designed the research,
845 analyzed the data, and wrote the article. All authors contributed to the discussion.

846

847

848 **Figure Legends**

849

850 **Figure 1.** Proteomic analysis of protein persulfidation in response to ABA in mesophyll
851 cells.

852 **(A)** Workflow of ABA leaf treatments followed by tag-switch protein labeling with CN-
853 biotin, protein purification, and quantitative SWATH analysis of eluted proteins.

854 **(B)** PCA representation plot of the 18 samples after SWATH analysis. Score for PC1
855 (70 %) versus PC2 (9.9%), Pareto scaling.

856 **(C)** Level of persulfidated ATG4a after ABA treatments. Values represent the mean
857 peak areas of extracted ion chromatogram of identified ATG4a peptides. *, p < 0.05.

858

859 **Figure 2.** Effect of sulfide on autophagy induced by ABA treatment.

860 **(A)** Immunoblot analysis of GFP-ATG8e fusion protein. One-week-old seedlings
861 expressing GFP-ATG8e fusion protein were transferred to liquid MS media and were
862 not treated (control) or treated with 50 µM ABA for 3 and 6 h and then 200 µM NaHS
863 for 1 h. Total protein extracts were prepared and subjected to SDS-PAGE and
864 immunoblot analysis with anti-GFP antibodies. Anti-tubulin antibodies were used as the
865 protein-loading control.

866 **(B)** Quantification of the free GFP/GFP-ATG8 ratio. For each condition, the levels of
867 free GFP and GFP-ATG8e fusion protein were quantified in a less exposed blot shown
868 in Supplemental Figure 6. The value of 100% was assigned to the free GFP/GFP-ATG8
869 ratio of the control sample. Data are from three independent experiments and evaluated
870 by two-factor ANOVA. Same letters indicate no significant differences. P < 0.05.

871

872 **Figure 3.** Persulfidation of *Arabidopsis* AtATG4a.

873 **(A)** Immunoblot analysis of persulfidated AtATG4a. Purified recombinant AtATG4a
874 was treated in the absence (-) or in the presence of 50 mM DTT (+) for 30 min at 4°C,
875 dialyzed, and subjected to the tag-switch labeling as described in the Methods. Then,
876 the proteins were subjected to immunoblot analysis using anti-biotin antibodies. Sypro
877 Ruby fluorescent staining is shown as the protein loading control.

878 **(B)** Analysis of AtATG4a using mass spectrometry. LC-MS/MS analysis of a tryptic
879 peptide of AtATG4a containing Cys170. The table inside the spectrum contains the
880 predicted ion types for the modified peptide, and the ions detected in the spectrum are
881 highlighted in red.

882 (C) AtATG4a protein sequence identified with 97% coverage. The peptide containing
883 persulfidated Cys170 is highlighted in yellow and all the Cys residues are red.

884

885 **Figure 4.** Arabidopsis ATG4a cleaves Chlamydomonas ATG8.

886 ATG4 activity of the recombinant AtATG4a was assayed by monitoring the cleavage of
887 CrATG8 from the unprocessed (CrATG8) to processed (pCrATG8) forms (indicated by
888 arrowheads) using SDS-PAGE followed by Coomassie Blue staining and quantification
889 of protein band intensities. The ATG4 activity (relative units) was determined as the
890 ratio of the band intensity of the processed CrATG8 to the sum of the intensities of the
891 unprocessed and processed CrATG8. Activity value of 1 corresponds to the maximum.
892 Processed pCrATG8 (lane 1) and unprocessed CrATG8 (lane 2) were loaded as
893 controls. Representative images are shown. Data are from three independent
894 experiments and evaluated by two-factor ANOVA. Same letters indicate no significant
895 differences. P < 0.05.

896 (A) Effect of incubation time. AtATG4a was incubated with CrATG8 in the absence or
897 in the presence of 10 mM DTT for the indicated times.

898 (B) Effect of DTT concentration. AtATG4a was incubated with CrATG8 in the absence
899 or in the presence of increasing concentrations of DTT for 4 h.

900 (C) Effect of TCEP concentration. AtATG4a was incubated with CrATG8 in the
901 absence or in the presence of increasing concentrations of TCEP for 4 h.

902

903 **Figure 5.** Effect of NaHS on AtATG4a enzyme activity.

904 (A) AtATG4a was incubated with 0.5 mM TCEP for 2 h and treated in the absence or in
905 the presence of indicated concentrations of NaHS for 1 h. Then, CrATG8 was added to
906 the incubation mixture, and ATG4 activity was monitored after 4 h using Coomassie-
907 stained gels as described in the Methods. All procedures were performed at 25°C. Lane
908 1 and 2, unprocessed and processed CrATG8, respectively. Lane 3, AtATG4a incubated
909 with CrATG8 in the absence of TCEP and of NaHS. Lanes 4-8, TCEP-pretreated
910 reduced AtATG4a incubated with CrATG8 in the presence of increasing concentrations
911 of NaHS (from 0 to 1 mM). A representative image is shown.

912 (B) Quantification of ATG4 activity (relative units) determined as the ratio of the band
913 intensity of the processed CrATG8 to the sum of the intensities of the unprocessed and
914 processed CrATG8. A value of 1 corresponds to AtATG4a in the absence of NaHS
915 (lane 4).

916 (C) Quantification of the protein band intensity corresponding to the monomeric
917 AtATG4a form marked by a rectangle in A. A value of 100% corresponds to AtATG4a
918 in the absence of NaHS (lane 4).

919 Data are from three independent experiments and evaluated by two-factor ANOVA.

920 Same letters indicate no significant differences. P < 0.05.

921

922

923 **Figure 6.** Effect of polysulfides on AtATG4a enzyme activity.

924 (A) AtATG4a was incubated with 0.5 mM TCEP for 2 h and subsequently treated in the
925 absence or in the presence of indicated concentrations of Na₂S₄ for 1 h. Then, CrATG8
926 was added to the incubation mixture, and ATG4 activity was monitored after 4 h using
927 Coomassie-stained gels as described in Methods. All procedures were performed at 25
928 °C. A representative image is shown.

929 (B) Quantification of ATG4 activity (relative units) determined as the ratio of the band
930 intensity of the processed CrATG8 to the sum of the intensities of the unprocessed and
931 processed CrATG8. A value of 1 corresponds to AtATG4a treated in the absence of
932 Na₂S₄ (lane 1).

933 (C) Quantification of the protein band intensity corresponding to the monomeric
934 AtATG4a form marked by a rectangle in A. A value of 100% corresponds to AtATG4a
935 treated in the absence of Na₂S₄ (lane 1).

936 (D) Reversibility of the effect. AtATG4a was incubated with 0.25 mM TCEP for 2 h
937 (lane 1) and treated with 100 µM Na₂S₄ for 1 h (lane 2) and 1 mM TCEP for 1 h (lane
938 3). ATG4 activity was monitored using Coomassie-stained gels. The experiment was
939 performed at least three times and a representative image used for the quantification of
940 the activity is shown.

941 Data are from three independent experiments and evaluated by two-factor ANOVA.

942 Same letters indicate no significant differences. P < 0.05.

943

944

945 **Figure 7.** Predicted structure of the AtATG4a-AtATG8a complex.

946 (A) 3D modelling of the AtATG4a-AtATG8a complex based on the structure of the
947 HsAtg4B-LC3 protein complex (PDB ID: 2Z0E). The AtATG8 protein sequence
948 (Q8LEM4 in UniProtKB) corresponds to the splice variant 1. Surface representation of
949 the protein complex and the equivalent residues surrounding the catalytic cavity

950 Cys170, Trp192, Asp364, and His366 in AtATG4a (red) and Phe116, Gly117, Thr120,
951 and Ala122 in AtATG8a (blue) are shown in the structural models.

952 **(B)** Zoomed view of the putative conformation of the active site showing the spheres
953 corresponding to the position and distance (Å) of catalytic residues Cys170, Trp192,
954 Asp364, and His366 in AtATG4a.

955

956 **Figure 8.** Cleavage and conjugation of Chlamydomonas ATG8 by Arabidopsis
957 proteins.

958 **(A)** ATG4 proteolytic activity in wild-type Arabidopsis leaves. Arabidopsis protein
959 extracts prepared from leaves of wild-type seedlings grown for 11 days on MS medium
960 were incubated with CrATG8 or site-directed mutant G120A proteins at 25°C for the
961 indicated times, and ATG4 activity was monitored as the cleavage of the ATG8 forms
962 to the processed (pCrATG8) forms by immunoblotting analysis with anti-CrATG8.
963 Processed pCrATG8 (lane 1), unprocessed CrATG8 (lane 2), and site-directed mutant
964 G120A (lane 6) were loaded as controls.

965 **(B)** ATG4 proteolytic activity in the Arabidopsis *atg4ab* mutant. Arabidopsis protein
966 extracts were prepared from the leaves of wild-type and *atg4ab* double mutant seedlings
967 grown for 7 days on the MS medium and transferred to the same medium (+N) or to a
968 nitrogen-deficient medium (-N) for additional 4 days. The protein extracts were
969 incubated with CrATG8 at 25°C, and ATG4 activity was monitored after 0, 0.5, or 2 h
970 as indicated by immunoblotting analysis with anti-CrATG8.

971 The arrowheads show the unprocessed CrATG8 and processed pCrATG8 protein bands,
972 and the asterisks indicate faster-mobility protein bands. Ponceau staining is shown as
973 the protein loading control of the Arabidopsis extract.

974

975 **Figure 9.** Sulfide inhibits the endogenous proteolytic activity of Arabidopsis ATG4.

976 **(A)** Effect of polysulfides on endogenous enzyme activity of Arabidopsis ATG4.
977 Arabidopsis protein extracts (Ex) were prepared from the leaves of seedlings grown for
978 11 days on the MS medium. The extracts were treated in the absence (un-Ex) and in the
979 presence of 200 µM Na₂S₄ (Na₂S₄-Ex), or 20 mM iodoacetamide (IAM-Ex) for 30 min,
980 or in the presence of 200 µM Na₂S₄ for 30 min and 1 mM TCEP for 30 min (Na₂S₄-Ex
981 + TCEP). Then, CrATG8 was added to the incubation mixture, and ATG4 proteolytic
982 activity was monitored. Lane 1, unprocessed CrATG8.

983 (B) Sulfide reverts the endogenous enzyme activity of Arabidopsis ATG4 under
984 autophagy-induced conditions. Arabidopsis protein extracts were prepared from the
985 leaves of seedlings grown for 7 days on the MS medium and transferred to the same
986 medium (+N) or to a nitrogen-deficient medium (-N) (left panel); or transferred to the
987 same medium (-mannitol) or to same medium containing 300 mM mannitol (+mannitol)
988 (right panel) for additional 4 days. The extracts were treated in the absence (un-Ex) or
989 in the presence of 200 μ M Na₂S₄ for 30 min (Na₂S₄-EX); CrATG8 was added to the
990 incubation mixture and ATG4 activity was monitored. Lane 1, processed pCrATG8 and
991 lane 2, unprocessed CrATG8.

992 The ATG4 activity was monitored at the indicated times by immunoblotting analysis
993 with anti-CrATG8. All procedures were performed at 25°C. Ponceau staining is shown
994 as the protein loading control.

995

996 **Figure 10.** Graphical model of ABA-triggered induction of autophagy mediated by
997 posttranslational modification of ATG4. Under basal conditions, intracellular sulfide
998 maintains high levels of persulfidation of the ATG4 pool, which inhibits the proteolytic
999 activity of the enzyme for ATG8 C-terminal processing. An increase in the intracellular
1000 level of ABA transiently decreases the level of persulfidation of the ATG4 population
1001 activating the protease activity of the enzyme and the processing of ATG8 that can be
1002 further lipidated to progress autophagy. Yellow circles represent ATG8 protein with or
1003 without the processed C-terminus (represented as X). Blue semicircles represent
1004 persulfidated ATG4 at the thiol group of Cys170 residue. Blue Pacman symbols
1005 represent ATG4 with reduced thiol group of Cys170 residue. The conjugation process
1006 of ATG8 with phosphatidylethanolamine (PE) and autophagosome initiation and
1007 closure are also shown.

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1009

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