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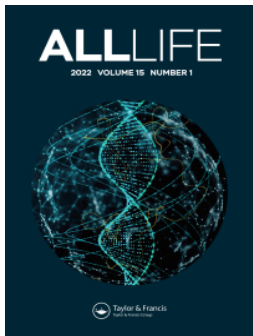
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Alternative splicing regulates autophagy in response to environmental stresses in cucumber (*Cucumis sativus*)

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ABSTRACT

Autophagy is important for cell survival during stress and nutrient recycle. However, the mechanisms involved in regulating autophagy in cucumber have not been determined. Here, we analyzed the transcript abundance of key autophagy genes in the conjugation pathway including *ATG3*, *ATG4*, *ATG5*, *ATG7*, *ATG8*, *ATG10*, *ATG12* and *ATG16* in response to nitrogen deficiency, pathogen infection and oxidative stress in *Cucumis sativus*. *ATG8* protein abundance was analyzed using immunoblot analysis. Seven out of 13 autophagy genes studied have splice variances including *ATG3*, *ATG4*, *ATG5*, *ATG8b*, *ATG8c*, *ATG8e* and *ATG10*. All except *ATG5* have changes in transcript variances abundance upon nitrogen starvation, indicating regulation via alternative splicing is pervasive in autophagy genes. This is the first report of alternative splicing in autophagy genes in cucumbers. However, transcription control is also important, especially in *ATG4*, and those without transcript variances. Moreover, different environment cues regulate autophagy genes through different mechanisms. The *ATG8* protein tag was also regulated at the protein level through post-translational modification and blockage of degradation. This work shows that the regulation of autophagy in cucumber is complex and involves many mechanisms. Better understanding of autophagy regulation would thus help breeding plants that perform better even under environmental stresses.

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Introduction

Autophagy, a nonspecific degradation and recycling of intracellular components, is important for cell survival, nutrient recycling and nutrient remobilization in various organisms (Mizushima 2007). In plants, it is particularly essential for mitigating the effects of stresses and environmental challenges (Liu and Bassham 2012). Various abiotic stresses including salt, drought, and hypoxia result in oxidative stress in plants. Autophagy is then induced to degrade proteins and cellular compartments damaged by the stress. Autophagy also plays an important role in the plant immune system to restrict pathogen invasion via hypersensitive response. Defects in the autophagy pathway cause plant to be sensitive to environmental stress, both biotic and abiotic, confirming the role of this process in stress responses (for a detail description of autophagy and plant stress response, see Signorelli

et al. 2019 and Su et al. 2020). The transcriptional control of genes involved in the autophagy process in responses to salt, drought and dark treatments, as well as phytohormones was confirmed in various plants (Xia et al. 2011; Pei et al. 2014; Wang et al. 2020a). Given the role of autophagy under stress conditions, understanding the regulation of this process could provide a way to improve crop adaptation and tolerance against environmental challenges (Liu et al. 2009; Sun et al. 2018a; Huo et al. 2020).

Autophagy is mediated by AuTophagy-related or ATG proteins. The cascade of activations leading to the autophagosome has been reviewed by Yang and Klionsky (2009) and involves several elements. *ATG12* is activated by *ATG7* before transferred to *ATG10*, which catalyzes the conjugation of *ATG12* to *ATG5*. The C-terminus of *ATG8* is removed by *ATG4*, a protease. Processed *ATG8* is then activated by *ATG7* and

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conjugated to phosphatidylethanolamine (PE) lipid by ATG3 and ATG12-ATG5-ATG16 complex. ATG8-PE is then integrated into membranes to subsequently form autophagosomes (for a detailed description of these molecular mechanisms, see Yang and Klionsky (2009)).

Alternative Splicing (AS) is a process in which more than one form of mRNA was produced as different splice sites were used. Up to 90% and 70% of all genes in human and Arabidopsis have splice variances (Pan et al. 2008; Calixto et al. 2018). This process alters the proteome landscape through changes in protein isoforms and quantity as well as introduces premature stop codons that induce nuclear-mediated decay and cis-regulatory elements targeted by miRNA and other regulatory machineries (Kazan 2003; Filichkin and Mockler 2012). In plants, splice variances were detected in association with differences in tissues, developmental stages and play an important role in plant response to environmental signals and stresses (Capovilla et al. 2018; Sun et al. 2018b; Zheng et al. 2019; Gao et al. 2021; John et al. 2021; Yan et al. 2021). Environmental stresses found to cause change in splicing in plants are pathogen infection, herbivore attack, extreme temperature, salinity, hypoxia, drought and deficiency of micronutrients (Laloum et al. 2018; Ganie and Reddy 2021; Lam et al. 2022). Genes targeted for AS upon stresses often have regulatory functions including transcription factors, splicing factors and protein kinase (Palusa et al. 2007; Rigo et al. 2019; Punzo et al. 2020). For example, a transcript variant encoding non-functional transcription factor was alternatively spliced to produce functional DREB2-type transcription factors in response to drought in rice (Matsukura et al. 2010). AS causes changes in protein structures of animal autophagy genes including ATG5, ATG7 and ATG8 (Liu et al. 2013; Ouyang et al. 2013; Park et al. 2016). Splice variances in autophagy genes were also detected in many plants including Arabidopsis, maize, rice (Chung et al. 2009; Wang et al. 2020b). In addition, AS of ATG4 during grape berry development was recently reported (Ma et al. 2023). However, the extent of how AS regulates autophagy in plant responses to stresses is not yet well understood.

Cucumber (*Cucumis sativus*, Cucurbitaceae family) has been used as a model plant for research in various aspects of abiotic stresses, nutrient limitation and pathogen resistance (Wei et al. 2015; Xin et al. 2017;

Dai et al. 2022). With the Cucumber genome available (Huang et al. 2009), autophagy gene orthologs were previously identified and their expression assessed (Han et al. 2021). Extensive transcriptome sequencing also provided abundant transcript data that could be explored by studying the role of AS in the responses to abiotic and biotic stress conditions.

Since AS regulates stress responses, we studied the regulation of AS upon autophagy genes and autophagy activity to establish if these processes take part in the stress response in cucumber. This work investigated the roles of AS, transcription and protein modifications that might affect functions of autophagy in cucumber. Here, we focus on key autophagy genes in the conjugation pathway, including ATG3, ATG4, ATG5, ATG7, ATG8, ATG10, ATG12 and ATG16. The expression of these autophagic genes and their splice variances under nitrogen starvation, oxidative stress and pathogen infection was analyzed, as well as ATG8 protein level.

Materials and methods

Plant treatments

The nitrogen starvation experiment was performed using 5-day-old seedlings of *C. sativus* grown hydroponically in liquid 1/2 MS medium with nitrogen before switching to 1/2 MS medium with and without nitrogen for 2, 4 and 6 days (Oka et al. 2012). The nitrogen source lacking in 1/2 MS medium without nitrogen was 21 mM NH_4NO_3 and 19 mM KNO_3 with the supplement of 18 mM KCl. First true leaves were collected from seedlings before the treatment (day 0) as well as the other six time points (day 2 +N; day 2 -N; day 4 +N; day 4 -N; day 6 +N; day 6 -N) for subsequent analysis. Oxidative stress was conducted using 2-week-old seedlings grown on soil. These plants were sprayed once with 50 μM Methyl Viologen (MV) prepared in 0.05% (v/v) Tween 20 (Song et al. 2005), and the first true leaves were collected before treatment as well as at 3, 6, 9, 12 and 15 days after the treatment. For pathogen responses, seven-day-old seedlings of *C. sativus* grown on soil were inoculated with sporangia of the fungi causing downy mildew disease (*Pseudoperonospora cubensis*) with the concentration of 1×10^4 sporangia mL^{-1} . Cotyledons were collected before inoculation and at 6, 12, 24, 72 and 168 h after inoculation. For each experiment, three replicates were

collected for each time point before being frozen in liquid nitrogen and stored at -80°C .

RNA isolation and reverse transcription

RNA was isolated from cucumber samples using CTAB supplemented with 1% beta-mercaptoethanol and 4% sodium dodecyl sulfate and subsequent extraction using chloroform and trizol. The RNA was then precipitated using ethanol with 0.2 M NaCl. Total RNA was treated with DNaseI to remove genomic DNA. First-strand cDNA synthesis was performed using one microgram of RNA, 0.25 mM oligo (dT) primers and 2 mM dNTP (Sreerathree et al. 2022). The mixture was denatured for 5 min at 65°C before adding MMuLV Reverse Transcriptase (Biotech rabbit, Germany) and incubated for 1 h at 42°C .

Quantitative real-time PCR

The real-time PCR analysis was performed in duplicate assays in 96-well plates; each 10 μl reaction consisted of $2\times$ QPCR Green Master Mix (Biotech rabbit, Germany), 5 μmol forward and reverse primers and 1 ng cDNA. The primers were designed to span the junction that differs between transcript variances and the sequences of primer are shown in Table 1. The data of Ct values obtained from two different experimental RNA samples were directly normalized to a house-keeping gene, *CsActin*. The fold-change in the expression of the gene of interest between each time point was then calculated as $2^{(-\Delta\Delta\text{Ct})}$. Statistical differences in the level expression compared to the expression at time point 0 or that of ATG7 were analyzed with student's *t*-test with two-tailed distribution.

Protein extraction and immunoblot analysis

Samples were ground and resuspended in extraction buffer (50 mM Tris-HCl pH 8.0, 150 mM NaCl, 2 mM EDTA, 10% (v/v) glycerol, 0.5% (v/v) IGEPAL CA-360, 1 mM phenylmethylsulfonyl fluoride) on ice. Crude proteins were separated on 18% SDS-PAGE before stained using Coomassie blue or transferred onto a PVDF membrane (GE Healthcare). Immunodetection of ATG8 was performed using anti-CrATG8 (Pérez-Pérez et al. 2010), with a working concentration of the primary antibody at 1:1000 in PBS containing 2.5% BSA. The Coomassie Brilliant Blue staining of

Table 1. Primers used in real-time RT-PCR.

Gene	Forward (5' to 3')	Reverse (5' to 3')
<i>CsATG5X1</i>	CCTTGTAAGGTGAAGATAG	CAGAACGCCATAACTCCAAC
<i>CsATG5X2</i>	GGAATCTAACGCGGATTAC	CTATATCCATCTCGCCAGTAC
<i>CsATG7</i>	AGCTTCTGAGCAGCCTCTTG	AAATCCATCCCCCTATTTTCG
<i>CsATG8aX1</i>	CAATCATTCGATTCCGTTGC	TCAGCCCTTCCACAATCACC
<i>CsATG8aX2</i>	GGGAGCTTTTCTTCCATGG	TCAGCCCTTCCACAATCACC
<i>CsATG8b</i>	GCATCAGGAGAAATACCC	ACAAACTGCCCAACAGTCAG
<i>CsATG8c</i>	CTGCTGACCTGACTGTTGG	ATGGCAGACATCAGTGATCC
<i>CsATG8dX1</i>	CTCTCTTAGTTTCAAGACTG	CATAAACGAACTGTCCACG
<i>CsATG8dX2</i>	CGCTCGTGGATTGTAGTTTC	CATAAACGAACTGTCCACG
<i>CsATG8eX1</i>	CTGGGGTTCTAACGTTTCGC	GCAATGTCACTCTCCACG
<i>CsATG8eX2</i>	GATCTTCTCTGCCACCAATC	GCAATGTCACTCTCCACG
<i>CsATG8eX3</i>	CTTCTCTGCGTCCGACCACC	GCAATGTCACTCTCCACG
<i>CsATG8eX4</i>	GTGGAAAGAGCCACCAATC	GCAATGTCACTCTCCACG
<i>CsATG8f</i>	TGGGAAATGTGAATCTCTG	CCTGGTGCTTAGAACATGG
<i>CsATG12</i>	GTCCATCCAGTTCTGCTCG	TGACAAACACAAACAAGGTGC
<i>CsPPH</i>	TGGGAAAGATTCTTCTGGG	TGCCCACAATGTAGACTGGT
<i>CsAPX</i>	TCCACCCTGGTAGAGAGGAC	AATGTCTGGTCCGAAAGAC
<i>CsRST</i>	TGCATTGCAAAGGACAATCG	GAAGTGTCCCTTGAGGTTTC
<i>CsActin</i>	ATTGTTCTCAGTGGTGGTTC	CCTTGAGATCCACATCTGC

the gel was used to estimate the amount of protein loaded.

Results

Splice variance in *CsATG* genes and its effects on the transcripts and protein-coding sequences

The splice variances of each ATG gene in the conjugation pathway were identified based on the Cucumber and NCBI databases. Only the splice variances detected by RT-PCR under any of the three conditions tested (nitrogen starvation, pathogen response and oxidative stress) were listed in Table 2, with the differences among splice variance described in detail as well as the condition that the splice variance was not detected. Out of 13 ATG genes assessed, seven genes contain multiple splice variances, including four variances for ATG3 and ATG8e, two variances for ATG4, ATG5, ATG8b and ATG8c and five variances for ATG10. Among these, only ATG4 and ATG5 have splice variances that affect the coding regions: N154H for *CsATG4* \times 1 and deletion at 98–127 amino acid position in *CsATG5* \times 2. The deletion is in the middle of conserved domain with Pfam ID of APG5 PF04106 and likely to affect the protein function. ATG3 variances alter 3'UTR after the stop codon, while the other four genes have splice variances that differ in 5'UTR with different forms of deletion. Because the splice variances of *CsATG3* are clustered within 20 bp after the stop codons, this region of *CsATG3* might have a regulatory function. That more than half of the genes in the cucumber conjugation pathway have multiple

Table 2. Summary of transcript variances confirmed in this work.

Gene	Gene ID	Cucumber ortholog	Splice variance	Accession number	Number of base pairs (bp)	Molecular weight of protein encoded (kDa)	Note
ATG3	LOC101210427	CsATG3	X1	XM_011658542.1	1486	35.492	108-bp deletion in 3'UTR at position 1126, 17 bp after the stop codon. No effects on protein sequence
			X2	XR_969726.1	1594	35.492	Full-length form
			X3	XM_004140463.2	1471	35.492	123-bp deletion in 3'UTR at position 1111, 2 bp after the stop codon. No effects on protein sequence
			X4	XR_969727.1	1579	35.492	15-bp deletion in 3'UTR at position 1112, 3 bp after the stop codon. No effects on protein sequence
ATG4	LOC101211819	CsATG4	X1	XM_011650784.1	1843	53.966	Two bases changed from TC in x2 to CA at positions +429 and +430, causing amino acid change at position 154 from H to N
ATG5	LOC101205497	CsATG5	X2	XM_004138902.2	1844	53.989	Higher expressed form in seedlings
			X1	XM_004140767.2	1331	39.552	Full-length form
			X2	XM_011652919.1	1243	36.007	30-amino acid deletion from positions 98 to 127, in PF04106 domain, 23 amino acid from the beginning of the domain and 2-bp addition at 5'end of the 5'UTR
ATG7	LOC101215499	CsATG7	–	XM_011659945.1	2326	77.715	–
ATG8	LOC101215846	CsATG8a	–	XM_011654340.1	835	13.520	–
	LOC101217161	CsATG8b	X1	XM_004151844.2	699	13.601	5-bp deletion from the position –178 to –174 and 88-bp deletion from the position –103 to –17 in the 5'UTR. No effects on protein sequence
	LOC101205863	CsATG8c	X2	XM_011651696.1	791	13.601	Full-length form
			X1	XM_004152360.2	927	13.559	14-bp deletion from the position –237 to –224 and a 24-bp deletion from the position –38 to –13 in the 5'UTR. No effects on protein sequence.
	LOC101204045	CsATG8d	X2	XM_004152359.2	965	13.559	Full-length form
			–	XM_011658569.1	873	13.373	–
	LOC101219454	CsATG8e	X1	XM_004134716.2	889	13.591	Full-length form
			X2	XM_004134717.2	759	13.591	130-bp deletion from the position –171 to –40 in the 5'UTR. Not detected in cotyledon of 7 and 8 d-old seedlings grown on soil used for the pathogen response experiment
			X3	XM_011659844.1	764	13.591	125-bp deletion from the position –166 to –40 in the 5'UTR
			X4	XM_011659845.1	660	13.591	Replacing the 270-bp region from the position –311 to –41 in the 5'UTR with a different exon (42 bp). Not detected in cotyledon of 7 and 8 d-old seedlings grown on soil used for the pathogen response experiment
	LOC101213890	CsATG8f	–	XM_004135913.2	583	13.514	–
ATG10	LOC101213064	CsATG10	X1	XM_011660912.1	1581	27.220	Replacing the region before the position –519 with a different exon to produce the 58-bp distal region of the 5'UTR. No effects on protein sequence
			X2	XM_011660913.1	1696	27.220	Full-length form
			X3	XM_004149270.2	1575	27.220	Replacing the region before the position –519 with a different exon to produce the 58-bp distal region of the 5'UTR, along with 6-bp deletion from the position –372 to –366. No effects on protein sequence.
			X4	XM_011660914.1	1550	27.220	145-bp deletion in 5'UTR from the position –518 to –373. No effects on protein sequence.
			X5	XM_011660915.1	1544	27.220	151-bp deletion in 5'UTR from the position –517 to –366. No effects on protein sequence.
ATG12	LOC101214589	CsATG12	–	XM_004150964.2	748	10.409	–
ATG16	LOC101219308	CsATG16	–	XM_011654969.1	1881	56.064	–

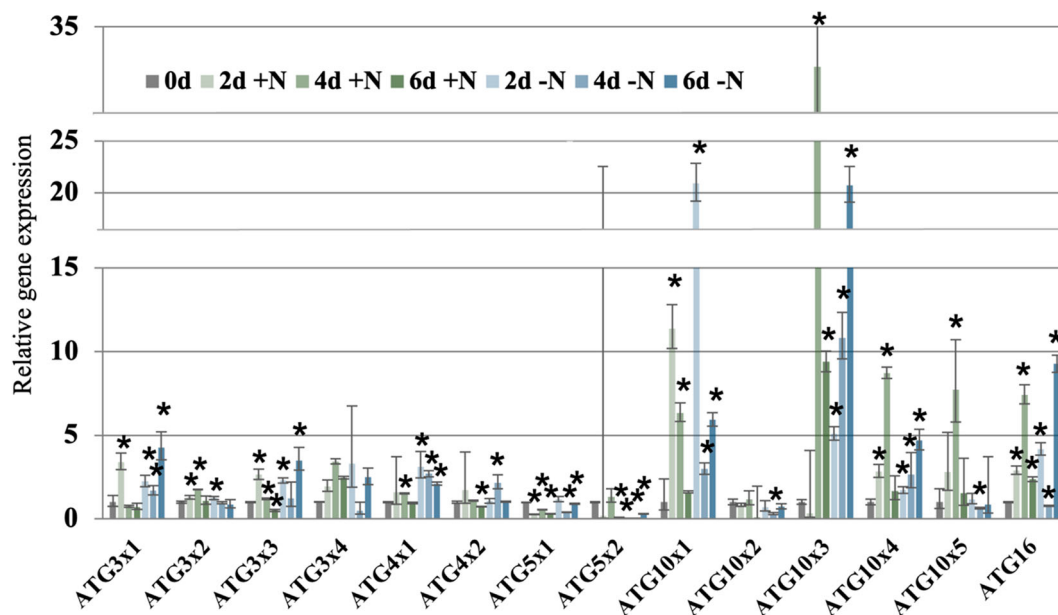


Figure 1. Relative expression of *ATG3*, *ATG4*, *ATG5*, *ATG10* and *ATG16* genes and their splice variances under nitrogen starvation. * indicates statistically significant difference (p -value less than 0.05) from the expression at 0 h based on Student's t -test.

splice variances suggests the importance of AS in the autophagy regulation. In addition, various modes of regulation are likely to be displayed as the AS observed in these genes affects protein isoforms and UTRs. This result also suggests the presence of cis-acting elements in the UTRs that could affect mRNA stability or translation efficiency.

Effects of environmental stresses on the abundance of the ATG splice variances

To understand the effects of environmental stress on the regulation of autophagy, we assessed the expression and the transcript variances of these *ATG* genes in seedlings under nitrogen starvation, and further investigated the key components (*ATG7*, *ATG8* and *ATG12*) under pathogen infection and oxidative stress. The expression level in the control of all *ATG8* genes was normalized to that of *ATG7* to determine the most abundant isoforms, with *ATG8f* consistently has the highest level of expression (data not shown). Changes in transcript abundance upon treatments indicate the regulation through transcriptional control. For genes with multiple splice variances, different patterns of responses among splice variances suggest the role of AS in regulating the response to stresses.

Nitrogen starvation increased transcript abundance for all *ATG* genes without splice variances including *ATG16*, *ATG7*, *ATG8a*, *ATG8d*, *ATG8f* and *ATG12*,

indicating the prominent role of transcription control on autophagy (Figure 1 and Figure 2A). For *ATG* genes with multiple splice variances, all except *ATG5* had different patterns of responses among splice variances. For example, only *ATG3 × 1* and *ATG3 × 3* abundance increased upon N starvation, but not the other two variances. In addition, a significant increase in abundance was detected for *ATG10 × 1*, *ATG10 × 3* and *ATG10 × 4* in all 3 samples under starvation, whereas *ATG10 × 2* and *ATG10 × 5* abundance decreased on day 4. The difference in response among splice variances of *ATG8* is also detectable, as the increase in *ATG8bx1* and *ATG8cx1* were more prominent than their splice variances, and *ATG8ex1* abundance significantly decreased upon starvation while the abundance of its variances increased. These results illustrate the pervasiveness of AS in the regulation of autophagy under nutrient stress.

Downey mildew inoculation did not change the expression of *ATG7*, *ATG8cx1*, *ATG8ex1* and *ATG8ex3* (Figure 2B). However, both variances of *ATG8b* had increases in expression by 6 h before reducing to levels before inoculation by 12 h. Furthermore, *ATG8a*, *ATG8cx2*, *ATG8d*, *ATG8f* and *ATG12* show a reduction in gene expression at 24 h after inoculation. This result indicates that the *ATG8b* might function as an early response to the pathogen infection, and this infection could block the accumulation of *ATG* transcripts potentially by inhibiting transcriptions or

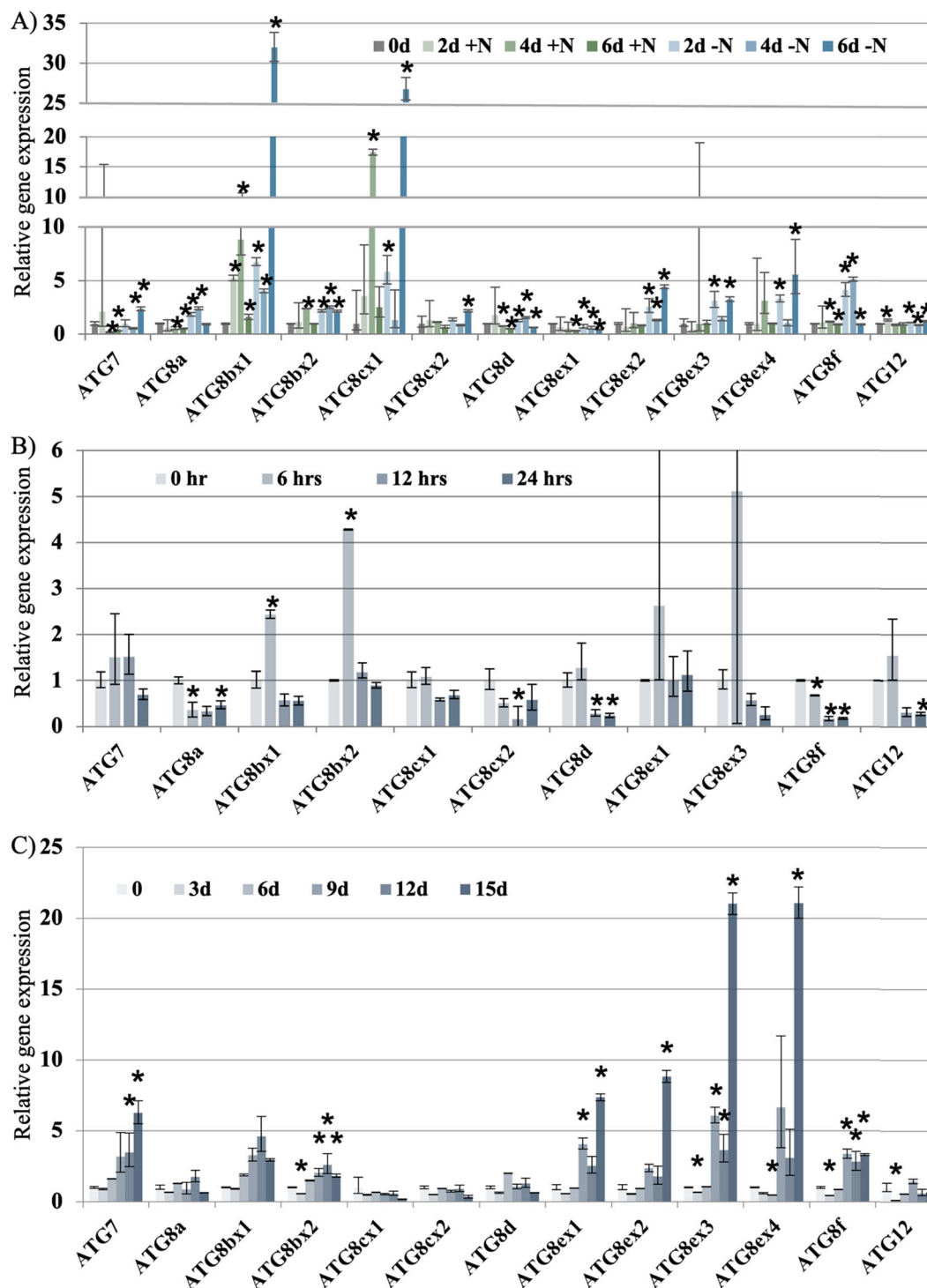


Figure 2. Relative expression of *ATG7*, *ATG8* and *ATG12* genes and their splice variants under nitrogen starvation (A), pathogen response (B) and oxidative stress (C). * indicates statistically significant difference (p -value less than 0.05) from the expression at Day 0 based on Student's t -test.

promoting transcript degradation. As a significance reduction in abundance among *ATG8c* transcripts was only detected for *ATG8cx2*, AS might also play a role in response to pathogen response.

Upon MV treatment, among the *ATG* genes without splice variants including *ATG7*, *ATG8a*, *ATG8d*, *ATG8f* and *ATG12*, only *ATG7* and *ATG8f* increase in expression especially in days 12 and 15 (Figure 2C).

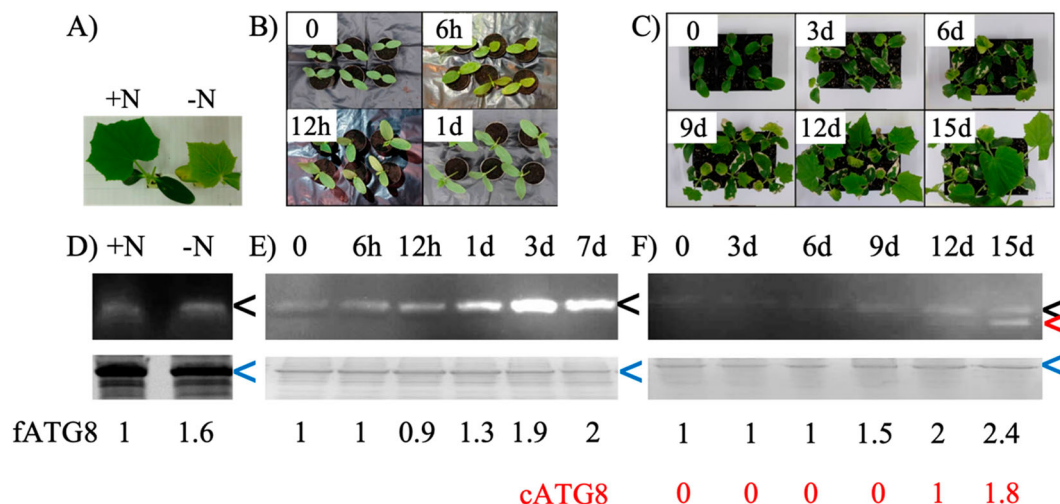


Figure 3. Phenotype of cucumber and immunoblot analysis of ATG8 protein abundance under nitrogen starvation (A, D), pathogen response (B, E) and oxidative stress (C, F). CsATG8 was detected using antibodies specific to CrATG8 (upper panel). Coomassie staining was used as a loading control (lower panel). Black, red and blue arrowheads indicate free CsATG8, conjugated CsATG8 and the band used for estimating the amount of total protein loaded, respectively. The numbers indicate the abundance of CsATG8 normalized to a loading control compared to control.

In addition, in this group, a significance reduction in expression at 3 days after treatment was detected for *ATG8f* and *ATG12*. For *ATG* genes with splice variances including *ATG8b*, *ATG8c* and *ATG8e*, there was no significance changes in expression detected for both variances of *ATG8c*, indicating a lack of either transcriptional or splicing control for this gene under oxidative stress. However, different response patterns among splice variances were detected in *ATG8b* as only *ATG8bx2* abundance was increased, but not *ATGbx1*. In addition, *ATG8ex1* and *ATG8ex3* increase in abundance happened earlier than the other variances. Therefore, the regulation of *ATG* expression was mediated through both transcription and AS. Moreover, *ATG8f*, *ATG8bx2*, and all four splice variances of *ATG8e* showed the highest transcript abundance increase under oxidative stress, suggesting that they are likely the main elements involved in the response to oxidative stress.

These results suggest that the transcriptional control as well as AS play a role in regulating *ATG* genes during abiotic and biotic stress response in cucumber, with different genes becoming targets of regulation by different mechanisms under each condition.

Effects of environmental stresses on ATG8 proteins

The level of ATG8 protein is often used as an indicator of autophagy levels since ATG8-PE conjugate is incorporated into the autophagosomal membrane

(Klionsky et al. 2008). Therefore, assessing the effects of environmental stresses on ATG8 proteins allows better understanding of their effects on autophagy level in general. Here, the abundance and the modification of cucumber ATG8 proteins under the stress conditions were assessed through western blot using rabbit anti-CrATG8 antibody. The predicted size of free CsATG8 (fATG8) is around 14 kDalton, whereas the conjugated form of CsATG8 (cATG8) appeared slightly smaller than the free form (Chung et al. 2009). Bands with the predicted size of fATG8 were detected in crude extracts from seedlings grown with and without nitrogen (Figure 3D and Figure 3A). However, once normalized to the Coomassie gel used as a loading control, there is a slight increase in ATG8 protein abundance from 1 to 1.6-fold under nitrogen starvation.

For the level of CsATG8 protein in response to pathogen infection, the sample for protein extraction was collected from both the early phase at 0, 6, 12, 24 h and the late phase at day 3 and day 7 after inoculation. Only one band at the predicted size of fATG8 was detected in all time points. The intensity of the band also increased along with time after inoculation, with a clear increase starting at 1 day (Figure 3E and Figure 3B). The ATG8 protein abundance increased from 1.3-fold after 1 day to 1.9-fold after 3 days and remained at a similar level after 7 days, suggesting that the accumulation occurred mostly between 3 and 7 days after inoculation. Upon treatment with MV, the band for

fATG8 was detected with increased intensity starting from 9 days after treatment (1.5, 2 and 2.4 folds in days 9, 12 and 15, respectively) (Figure 3F and Figure 3C). In addition, another band for cATG8 was detected starting from 12 days after treatment and increase in intensity to 1.8-fold in day 15.

The increase in ATG8 protein abundance during these abiotic and biotic stresses indicated the roles of ATG8 protein and autophagy in cucumber responses to stress. In addition, the detection of the smaller size band suggested a role of post-translation modification in oxidative stress response.

Discussion

Our work shows that AS is frequent in the autophagy response, with 7 out of 13 cucumber autophagy genes studied here having multiple splice variances. In animals, AS of autophagy genes was reported to change the coding regions that affect protein structures of ATG5, ATG7 and ATG8 (Liu et al. 2013; Ouyang et al. 2013; Park et al. 2016). Here, AS of *ATG5* allows changes in the coding sequence that potentially alter the protein function. However, as both splice variances of *ATG5* have the same pattern of changes under N starvation, *ATG5* was likely regulated mainly through transcriptional control under nutrient stress. In some cases, the regulations through the transcription control or AS could be selective depending on the stress conditions. AS affects *ATG8b* and *ATG8e* response to nitrogen starvation and oxidative stress but not during pathogen infection, whereas regulation through AS of *ATG8c* occurs during nitrogen starvation and pathogen infection but not under oxidative stress. In addition, for the *ATG* genes with splice variance, their expression was mainly regulated through both AS and transcription concurrently. These results illustrate the complex role of AS in regulated specific responses to stresses in plants (Wang et al. 2020b; Chaudhary and Kalkal 2021).

Transcription control is also important for cucumber response to abiotic and biotic stresses. Indeed, *ATG8f*, one of the genes with no transcript variance, has the highest expression level in the control group for all three experiments. Even though such high expression level might be partly due to the differences in the amplification efficiency of primers used, the increases of *ATG8f* expression in response to N starvation and oxidative stress suggest that *ATG8f* could

be one of the key autophagy genes regulated by transcriptional controls during environmental stresses in cucumber. In rice, *OsATG8a* is the most abundant transcript and *OsATG8d* the most strongly induced upon N starvation and abiotic stress (Xia et al. 2011). The differences among the most abundant ATG8 isoforms and their role in response to environmental cues in different plant species illustrate the evolutionary divergence in the function of this gene. The reduction in *ATG8f* transcript abundance within a day after Downey mildew inoculation also indicates its significance as a target of transcription suppression in response to the pathogen. Wheat *ATG8* also shows reduced expression upon pathogen infection in susceptible plants but is upregulated in resistant plants (Zhang et al. 2020). The expression of *ATG12*, another protein tag for the conjugative pathway, did not change significantly under the three treatments presented here. Given that *ATG12* expression was previously shown to increase upon N starvation in rice and maize (Chung et al. 2009; Xia et al. 2011), this suggests a minor role of *ATG12* under stress conditions in cucumber. In cucumber, *ATG7* was responsive to abiotic stress, but not to biotic stress. *ATG7* expression increases in maize under N starvation and carbon starvation, in *Arabidopsis* under heat treatment and sucrose starvation and in rice under salt treatment and nitrogen starvation (Rose et al. 2006; Chung et al. 2009; Xia et al. 2011; Zhou et al. 2013). These data indicate that *ATG7* is transcriptionally regulated by abiotic stress across species.

An additional layer of control in autophagy is at the protein level. The increase in ATG8 protein abundance is consistent with the increases of transcripts of *ATG8* gene family during N starvation, oxidative stress and pathogen infection, and this is quite conserved among plant species (Chung et al. 2009; Pérez-Pérez et al. 2010; Pérez-Martín et al. 2014). *ATG8a* and *ATG8f* are potentially responsible for the increases in ATG8 proteins during N starvation, and *ATG8b*, *ATG8e* and *ATG8f* are during oxidative stress. The selective induction of different *ATG8* isoforms under different conditions was also observed in *Arabidopsis*, maize, rice and wheat (Yoshimoto et al. 2004; Chung et al. 2009; Xia et al. 2011; Pei et al. 2014). In contrast, upon pathogen infection, only *ATG8b* transcript increased, whereas the rest of the *ATG8* family reduced in expression after only one day of inoculation. The increases in *ATG8b* transcript abundance seemed too low to be accounted

for the increased protein level. This increase in ATG8 protein in contrast to the little increase in *ATG8* transcripts may indicate either increase in translation or delay in protein degradation as a result of pathogen infection.

Post-translation modification of ATG8 through the conjugation of ATG8 onto PE before its integration into autophagosomal membrane is critical for ATG8 function (Yang and Klionsky 2009). In cucumber, a smaller band representing the PE-conjugated form of ATG8 became detectable after 12 days of oxidative stress confirming the modification of ATG8 and its role in oxidative responses, similar to reports in other organisms (Chung et al. 2009; Suttangkakul et al. 2011). In addition, the increase in transcript abundance of *ATG7*, the gene critical for the ATG8-PE conjugation, in days 12 and 15 also correlates with the detection of the PE-conjugated form of ATG8. Such increases in transcript and protein abundance in days 12 and 15 after MV treatment could also be due to a developmental process, namely senescence, since these first true leave samples were collected from 26- and 29-day-old plants at the age when cotyledon senescence can be detected (Kim 2004). ATG8-PE conjugate accumulation in cucumber in response to senescence is consistent with that detected in leaf senescence in maize (Chung et al. 2009). No detection of the conjugated form in N starvation and pathogen infection could imply a high turnover rate of the conjugated form or the lack of conjugation activity under these conditions. For N starvation, the first scenario is more likely as the conjugated form is normally present in the N starvation condition as shown in *Arabidopsis*, maize and *Chlamydomonas* (Chung et al. 2009; Pérez-Pérez et al. 2010; Suttangkakul et al. 2011). In pathogen infection, lack of conjugation could also be the case, based on the decrease in transcript abundance of most *ATG8* genes. The targets of gene downregulation for *Pseudoperonospora* appeared to be *ATG8s* and *ATG12*, which encode protein tags, but not *ATG7* (Zhang et al. 2020).

This work shows the regulation of autophagy genes in the conjugation pathway in cucumber under three different stress conditions. In addition to the transcriptional control, the autophagy process was also regulated post-transcriptionally both through AS and at the protein level, which can affect protein function as well as protein degradation. This work is also the first detailed analysis of splice variances in autophagy genes

in plants. Further work will establish the exact mechanisms of protein accumulation and cis-regulatory elements in the 5' and 3'UTRs that can deepen our understanding of cucumber tolerance under environmental stresses and suboptimal conditions.

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Disclosure statement

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Author contributions

AS, PT and SV conceived the project and designed experiments. PT, TC, WP and PN performed the experiments. PT, PN, SV and AS analyzed the data. PT, SV, LG and AS interpreted the data, prepared Figures and Tables and wrote the manuscript. All authors have approved the final version.

Data availability

The data that support the findings of this study are openly available in Mendeley Data: <http://doi.org/10.17632/v6f2rhv42z.2>.

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