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Vacuolar processing enzymes, AmVPE1 and AmVPE2, as potential executors of ethylene regulated programmed cell death in the lace plant (Aponogeton madagascariensis)

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Running title: Role of vacuolar processing enzymes in lace PCD

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Summary

Perforation formation in *Aponogeton madagascariensis* (lace plant) is an excellent model for studying developmentally regulated programmed cell death (PCD). In this study, we isolated and identified two lace plant vacuolar processing enzymes (VPEs) and investigated their involvement during PCD and throughout leaf development. Lace plant VPE transcript levels were determined during seven different stages of leaf development. PCD and non-PCD cells from “window” stage leaves (in which perforations are forming) were separated through laser-capture microscopy and their transcript levels were also determined. VPE activity was also studied between the cell types, through a VPE activity-based probe JOPD1. Additionally, VPE transcript levels were studied in plants treated with an ethylene biosynthesis inhibitor, aminoethoxyvinylglycine (AVG). The two isolated VPEs, *Am*VPE1 and *Am*VPE2, are vegetative type VPEs. *Am*VPE1 had higher transcript levels during a preperforation developmental stage, immediately prior to visible signs of PCD. *Am*VPE2 transcripts were higher later during window and late window stages. Both VPEs had higher transcript and activity levels in PCD compared to non-PCD cells. AVG treatment inhibited PCD and associated increases in VPE transcript levels. Our results suggested that VPEs are involved in the execution of the ethylene related PCD in the lace plant.

Keywords: *Aponogeton madagascariensis*, ethylene, leaf development, perforation formation, programmed cell death, vacuolar processing enzymes
**Introduction**

Programmed cell death (PCD) is regulated cell demise necessary for elimination of specific cells in most multicellular organisms, development, adaptation and response to stress (Greenberg 1996; Lam 2004). It has been observed in unicellular and multicellular organisms, including fungi, plants and animals (Rantong and Gunawardena 2015). In mammals, apoptosis (a form of PCD) is modulated and executed by caspases. Although the execution of PCD through caspases is well understood in animals, execution of PCD in plants is still unclear. In addition, plant species whose complete genomes have been sequenced lack genes encoding true caspases. However, some of the hallmarks of animal PCD, which are directly or indirectly effects of caspases, are also observed in plants. These include chromatin condensation, DNA laddering, release of cytochrome c, shrinkage of the cytoplasm, and activation of death proteases (Wang et al. 1996; Adrain and Martin 2001; reviewed in Elmore 2007; reviewed in Reape and McCabe 2008). This suggests the presence of proteins that perform caspase-like duties during plant PCD. Identification of such proteins led to the isolation of saspases (Dodson and Wlodawer 1998; Coffeen and Wolper 2004), phytaspases (Chichkova et al. 2010; Cai and Gallois 2015), metacaspases (Uren et al. 2000), and vacuolar processing enzymes (VPEs; Kinoshita et al. 1995a, b; reviewed in Bonneau et al. 2008; Cai and Gallois 2015).

VPEs share many characteristics with animal caspases. They are both cysteine proteases, have a His-Cys catalytic dyad, are synthesized as inactive proenzymes, can cleave after Asp residues in similar synthetic substrates and share similar inhibitors (Hatsugai et al. 2004; Rojo et al. 2004; Misas-Villamil et al. 2013). VPEs are more similar to caspase-1 in terms of their catalytic dyad, inhibitors and substrates (Wilson et al. 1994; Cohen 1997; Hiraiwa et al. 1999; Nicholson, 1999; Hara-Nishimura et al. 2005; Sanmartin et al. 2005). Due to the similarities with caspases, VPEs have been isolated in several species and their role in plant PCD is under investigation. In tobacco, caspase-1 activity required for PCD to occur during hypersensitive (HR) in response to TMV is attributed to VPEs (del Pozo and Lam 1998; Hatsugai et al. 2004). VPEs also play an important role in other types of plant PCD such as embryogenesis (Hara-Nishimura et al. 2005), seed development (Nakaune et al. 2005), and leaf senescence (Kinoshita et al. 1999). In *Arabidopsis*, a total of 4 VPEs have been isolated and divided into two types (Kinoshita 2009).
et al. 1995a, b; Nakaune et al. 2005; Yamada et al. 2005). Seed type VPEs, delta-VPE and beta-VPE, are mostly expressed in seeds. Vegetative type VPEs, alpha-VPE and gamma-VPE, are mainly expressed in vegetative tissue.

Lace plant (*Aponogeton madagascariensis*) is a submerged aquatic monocot which forms perforations in its leaves during normal development by way of PCD (Fig. 1; Gunawardena et al. 2004). Lace plant is one of the few plant species known to employ PCD during leaf morphogenesis. However, unlike in the other plant species such as *Monstera obliqua*, lace plant PCD regions are easily visible and occur at highly predictable locations (Figs. 1B-E). Lace plants can also be propagated in microbe-free conditions in magenta boxes for experimental purposes (Gunawardena et al. 2006; Fig. 1B). In addition, the lace plant is ideal for light microscopy due to its thin and transparent leaves. Lace plant leaf development has previously been divided into seven stages (Rantong et al. 2015). These stages are early preperforation, preperforation, early window, window, late window, mature and senescence stage. Visible signs of perforation-related PCD are observable during the early window, window and late window stages.

Ethylene plays a role in lace plant leaf development via PCD during perforation formation (Dauphinee et al. 2012; Rantong et al. 2015). In addition, a caspase-1 inhibitor (Ac-YVAD-CMK) is known to stops PCD and perforation formation in the lace plant, which provides indirect evidence for the involvement of VPEs in lace plant PCD during leaf morphogenesis (Lord et al. 2013). Due to the involvement of caspase-1 like activity and ethylene in lace plant PCD signaling, we investigated the interactions between VPEs and ethylene throughout perforation formation. Although links between ethylene and VPEs during PCD have been studied in the HR, to our knowledge no studies have investigated their relationship during developmentally regulated PCD.

The objective of this study was to isolate lace plant vegetative type VPEs and study their expression patterns and activity during developmentally regulated PCD and leaf development in the lace plant. Recently we showed that ethylene plays a role in lace plant PCD (Dauphinee et al. 2012; Rantong et al. 2015), therefore, we also investigated the
effect ethylene biosynthesis inhibition on VPE transcript levels during leaf development via developmentally regulated PCD.

**Materials and Methods**

**Plant propagation**

Lace plants were propagated as described by Gunawardena et al. (2006), under sterile conditions, and in Magenta GA7 boxes. Plants were kept in 12 h light/12 h dark cycles. The light intensity was approximately 125 µmol m\(^{-2}\) s\(^{-1}\) and was provided through daylight simulating fluorescent bulbs (Philips, Daylight Deluxe, F40T12/DX, Markham, Ontario, Canada). The plants were maintained at 24°C.

**RNA extraction and cDNA Synthesis**

RNA was extracted from leaves sampled from seven different developmental stages described in Rantong et al. (2015). These leaf developmental stages are early preperforation, preperforation, early window, window, late window, mature) and senescence. Four RNA samples were collected from each leaf developmental stage. In total, 28 independent RNA samples were analyzed. Each RNA sample consisted of tissue from at least 3 leaves from different plants. RNA quality was determined through gel electrophoresis and spectrometry at 260 nm. Synthesis of cDNA was performed as described in Rantong et al. (2015).

**PCR amplification of lace plant VPE cDNA**

Initial VPE fragments were amplified degenerate primers. The degenerate primers were designed using alignments of VPE sequences from different species, performed through CLC combined workbench (CLC Bio-Qiagen, Aarhus, Denmark). The sequences used in the alignment were from *Arabidopsis thaliana* (NM_128154), *Ricinus communis* (D17401), *Vitis vinifera* (XM_002276723), *Populus trichocarpa* (XM_006371798), *Populus tomentosa* (FJ461342), *Malus hupehensis* (FJ891065), *Solanum tuberosum* (NM_001288343), *Zea mays* (NM_001111649 and AJ131719), *Hordeum vulgare* (AM941114), *Beta vulgaris* (AJ309173), *Nicotiana tabacum* (AB075947 and AB075948), and *Solanum tuberosum* (EU605871). The forward and reverse degenerate primers used for amplification of the initial fragments of *AmVPE1* are 5’-

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TGGCYRTGCTCATYGCCGGCTC-3’ and 5’-ACCARGCTTTTDTAGGTYCC-3’, respectively. Forward (5’-TGGGCYGTCCTSMTCGCCGG-3’) and reverse (5’-GGCATCCCAAGRACHCCAGGTCC-3’) degenerate primers were used to amplify a fragment of AmVPE2. The PCR reactions consisted of 3.5 µl of cDNA, 11.15 µl nuclease free water, 1 µl of 10 mM dNTP mix, 1 µl of 10 mM forward and reverse primer, 2 µl 10X Thermobuffer, and 0.35 µl of 5 U/ µl Taq DNA polymerase. PCR conditions consisted of initial denaturing at 94ºC for 5 minutes, 40 cycles at 94ºC for 30 seconds (denaturing), primer annealing at 48ºC for 30 seconds, elongation at 72ºC for 1 minute, and final elongation at 72ºC for 10 minutes. After sequencing of the initial lace plant VPE fragments, 3’-RACE was used to isolate the rest of the 3’ end, including 3’-UTR using an abridged universal amplification primer (AUAP; 5’-GTACTAGTCGACGCGTGCC-3’) and an anchored primer (AP; 5’-GGCCACGCGTGACTAGTACTTTTTTTTTTTTTTTTTTTTTT-3’).

**Sequence analysis**

Cloning and sequencing was performed as described in Rantong et al. (2015). BioEdit Sequence Alignment Editor (Carlsbad, Ottawa, Ontario, Canada) was used to trim low quality portions and vector sequences. It was also used to deduce the amino acid sequences from nucleotide data. Both nucleotide and amino acid sequences were then compared with National Center for Biotechnology Information (NCBI) nucleotide collection (blastn) and nonredundant protein (blastx) database sequences, respectively. ClustalW2 (Larkin et al., 2007) was used to calculate sequence percentage identities for both nucleotide and amino acid sequences.

**Phylogenetic analysis**

The lace plant VPE sequences were aligned with 34 known VPE sequences from other plant species using CLUSTALW (Thompson et al. 1997). The VPE sequences used for phylogenetic analysis were from both monocot and dicot species. MEGA version 6.06 was used to construct a single tree using the Neighbor-Joining method. A gap at the 5’ in lace plant sequences and the corresponding amino acids in all other sequences was removed prior to construction of the phylogenetic tree. Branch strength was calculated
using 1000 replicates in a nonparametric bootstrap test. The rice putative asparagine-specific endopeptidase precursor (NP_910213) was used as an out-group.

**Analysis of VPE transcript levels throughout leaf development**

VPE transcript levels were measured in the seven different stages of lace plant leaf development. *AmVPE1* and *AmVPE2* transcripts were detected in cDNA from all leaf developmental stages. The primers used for quantitative PCR (QPCR) did not amplify any products from genomic DNA, but amplified expected-size products in cDNA samples. Actin was used as a reference gene. Transcript levels from four different RNA samples per leaf developmental stage (28 independent RNA samples in total) were analyzed. Transcript levels were determined through QPCR. The QPCR reactions were performed in a Rotor-Gene 3000 (Corbett Research, Sydney, Australia) using QuantiTect SYBR Green PCR Kit (Qiagen, Mississauga, Ontario, Canada), following the manufacturer’s instructions. The forward and reverse primers used for amplifying *AmVPE1* fragments are 5’-GCATTGTAAAGGAGCGGACA-3’ and 5’-TTGTACCCATAAACAAGGCAAT-3’ respectively. Forward primer 5’-CCGAGTGGTTAAAGAACGAA-3’ and reverse primer 5’-CTGAACCAATGTACAAGGCAAGTG-3’ were used to amplify fragment of *AmVPE2*. Prior to QPCR, these primers were used in regular PCR to amplify fragments. The fragments were ligated into a pGEM3T vector, cloned, sequenced and verified to be the expected fragment. The QPCR conditions used consisted of 95°C for 15 min (initial denaturing), 40 cycles of 94°C for 20 s (denaturing), primer annealing at 60°C for 20 s, and 72°C for 30 s (elongation). Product purity was determined by observing the melting temperature curve at the end of QPCR. A standard curve of copy number was generated from quantified target sequences for each gene. The standard curves were used to determine mRNA copy numbers as explained in Bustal et al. (2005). Absolute mRNA copy number of each VPE was divided by *AmActin* (KR779004.1) mRNA copy number to obtain a mean normalized expression value in each sample. Actin primers used for QPCR are 5’-TACGACAGGTATCGTGCTTG-3’ and 5’-CAAGCACGATACCTGTCGA-3’.
Analysis of VPE transcript levels between dying (PCD) and non-PCD (NPCD) cells

To further examine the role of VPEs in lace plant PCD, transcript levels were determined in PCD and NPCD cells. A Zeiss PALM Laser Capture Microdissection and Imaging System (North York, Ontario, Canada) was used to separate PCD and NPCD cells. The cells were collected from early window and window stage leaves, in which the cells are easily visually distinguishable through color differences. NPCD cells have high amounts of anthocyanin and are pink. Most of the anthocyanin is lost in PCD cells (Figure 1D). Cells were collected from at least three different leaves per sample and a total of 8 different samples (four samples per cell type) were used for RNA extraction. A ReliaPrep RNA Cell Miniprep kit (Promega, Nepean, Ontario, Canada) was used for RNA extraction, following manufacturer’s instructions. RNA was treated with DNase 1 prior to cDNA synthesis. A Protoscript M-MuLV First Strand cDNA Synthesis Kit (New England Biolabs, Pickering, Ontario, Canada) was used for cDNA synthesis.

Labelling of leaf tissue with VPE activity probe JOPD1

Labelling was performed using a protocol modified from Lu et al. (2015). Window stage lace plant leaves were dissected into sections of approximately 9 mm². The labelling was performed in 300 µl total volume containing distilled water and 0.5% silwett L77 (Shawnee Mission, Kansas, USA). For treatments, the leaf tissue was incubated in 4 µM JOPD1 probe solution (kindly donated by Dr. Renier van der Hoorn, Max Planck Institute for Plant Breeding Research, 50829 Cologne, Germany) for 24 h at room temperature under constant shaking. In no-probe-controls, an equal volume of DMSO was added. JOPD1 is an activity-based BOPIDY tagged probe that targets VPEs. Details on the synthesis and mode of action of JOPD1 are provided in Lu et al. (2015) and Ashnest et al. (2015). In short, a permanent covalent bond is produced between VPEs and JOPD1 when the active VPEs catalyze the cleavage of the probe. This interaction produces fluorescence detectable through a confocal microscope. A Nikon C1 Confocal Imaging System with an Eclipse Ti inverted Microscope (Nikon, Mississauga, Ontario, Canada) was used to detect the fluorescence of the BOPIDY fluorescence tag (excitation 561 nm/emission 595/50 nm). For each experiment, there were 4 replicates and the experiment was repeated three times.
AVG treatment
Plants were treated with 5 µmol/L AVG (Sigma Aldrich, St. Louis, Missouri, USA) as described in Dauphinee et al. (2012). Leaf tissue was harvested from plants three weeks post-treatment, and the plants had produced three-four new leaves. The leaves were separated into six stages of leaf development; early preperforation (EPP), preperforation (PP), early window (EW), window (W), late window (LW) and mature (M). Senescent stage leaves were not available in this experiment since the plants were harvested before any of the leaves reached senescence. RNA was extracted from leaves at each of the six developmental stages. At least three different leaves from different plants were used in each RNA sample. A minimum of four independent RNA samples per leaf developmental stage (24 independent samples in total) were used. cDNA synthesis and QPCR were also performed as described above.

Statistical analysis
GraphPad Prism version 5.00 (San Diego, California, USA) was used to analyze Quantitative PCR data. Relative abundance of gene transcripts is presented as mean ± S.E.M. For leaf developmental stages, significant differences in the number of transcripts were determined through a general linear model of variance (ANOVA). Means of individual treatments were compared using a Tukey post-test, if the overall relationships were significant. An unpaired t-test was used for comparisons of relative transcript levels between PCD and NPCD cells. For the AVG experiment, a two-way ANOVA with no repeated measures, followed by Bonferroni post-test, was used to determine significant differences in transcript levels. Data was determined to be statistically significant if \( P < 0.05 \).

Results
Lace plant VPEs
Two vegetative type lace plant VPEs, AmVPE1 (KR779002) and AmVPE2 (KR779003), were isolated. Their isolated fragments were 1436 and 1603 base pairs, respectively. The entire 3’ end of their cDNA was isolated, including the untranslated region. At the nucleotide level, AmVPE1 and AmVPE2 fragments were 83.6 % identical. They both translated into 437 amino acid fragments (Fig. 2A). At the amino acid level, they were
88.3 % identical. They were both more identical to *Arabidopsis* vegetative type VPEs than to the seed type. *AmVPE1* was 69 % and 71 % identical to *Arabidopsis* gamma and alpha-VPEs respectively. It was only 58 and 53 % identical to the seed type *Arabidopsis* VPEs (beta and delta-VPE, respectively). *AmVPE2* shared 68 % and 70 % identity to *Arabidopsis* gamma and alpha-VPEs respectively, while it was only 60 % and 52 % identical to the *Arabidopsis* beta and delta-VPEs, respectively.

The domains and sites identified within *AmVPE1* and 2 that are conserved in *Arabidopsis* vegetative VPEs include the mature protease domain, essential amino acids (Cys102, His191 and Cys241), N-glycosylation site (Asn360), and a C-terminal propeptide (Figure 2A). Most of these are also conserved in *Arabidopsis* seed-type VPEs, except His191, which is an essential amino acid, and is instead replaced by an Asn in delta-VPE. Also, less of the C-terminal propeptide region is conserved between lace plant VPEs and *Arabidopsis* seed-type VPEs, compared to vegetative type *Arabidopsis* VPEs (Fig. 2A).

Phylogenetic analysis comparing the amino acid sequence of *AmVPE1* and *AmVPE2* with 34 VPEs in other plant species, also revealed that they share more in common with the vegetative type VPEs than seed type VPEs (Fig. 2B). *AmVPE1* and 2 also appear to be more related to the monocot (rice and maize) vegetative type VPEs than the dicot vegetative VPEs (Fig. 2B).

**AmVPE1** and **AmVPE2** transcript expression levels in different stages of lace plant leaf development

Transcript level results demonstrated that the lace plant VPE transcripts are differentially expressed during leaf development. *AmVPE1* transcripts levels in leaves sampled at the preperforation stage were significantly higher than at all other stages of leaf development (Fig. 3A). This stage of leaf development is immediately prior to obvious visible signs of PCD and perforation formation. During the early window and window stage, the *AmVPE1* transcript levels were approximately 2-fold lower than in preperforation. During the late stages of PCD, in late window stage, the *AmVPE1* transcript levels declined to about 3-fold less than in the early window and window stages. In mature and senescence stage leaves, the *AmVPE1* transcripts are at their lowest levels as compared to all the other stages of leaf development. *AmVPE2* transcript levels were about 1.4-fold higher in
window and late window stage leaves (in which rapid cell death was occurring) than in other stages (Fig. 3B). There were no significant differences in AmVPE2 transcript levels among all the other stages (early preperforation, preperforation, early window, mature and senescence) of leaf development. AmVPE2 transcript levels were approximately 1000-fold higher than AmVPE1 transcript levels. AmActin transcripts were constitutively expressed throughout leaf development (Fig. 3C).

AmVPE1 and AmVPE1 transcript levels in PCD versus NPCD cells
QPCR showed that PCD cells had significantly higher transcript levels of both lace plant VPEs than the NPCD cells (Figs. 4A and B). AmVPE1 transcript levels were about 3.2-fold higher in PCD than in NPCD cells. The amount of AmVPE2 transcripts in PCD cells was more than double the amount in NPCD cells. AmVPE2 transcript levels appeared to be higher (1000-fold) than AmVPE1 transcript levels at the cell level as well. AmActin transcript levels were not significantly different between the two cell types (Fig. 4C).

VPE activity in PCD compared to NPCD cells
Probe-associated fluorescence (red) was observed within vacuoles, where VPEs are known to be active (Fig. 5). The activity-based probe is an inhibitor tagged with a reporter (BOPIDY fluorescent reporter) and reacts specifically with VPE active-site residues. The VPE activity and fluorescence was higher in PCD cells compared to NPCD cells. Dead cells (late PCD cells) did not display probe-associated fluorescence (Figs. 5E and F). No probe-associated fluorescence was detected in the negative controls (Fig. 5D).

Effects of AVG treatment
Leaf Morphology
Representative leaf layouts were assembled to highlight leaf morphological differences at the six stages of development and between treatments (Fig. 6). Early preperforation (EPP) stage leaves (Figs. 6A and B) had just emerged from a corm, were light green, tightly furled and had no signs of perforation formation and PCD. Preperforation (PP) leaves (Figs. 6A and B) were still furled, and also did not show obvious signs of perforation formation or PCD. At the early window (EW) stage, leaves in control plants started to show signs of perforation formation. The perforation sites started to become
somewhat transparent as PCD cells in these leaves started to lose their red pigmentation (Fig. 6A, leaf EW). In the AVG treated plants, however, the signs of perforation formation were minimal or nonexistent in early window stage leaves (Fig. 6B, leaf EW). The leaves in both the AVG treated and control plants were about half-unfurled at this stage. During the window (W) stage in control plants, perforation formation had progressed and leaves were actively forming perforations (Fig. 6A). In the AVG treated plants however, most of the window stage leaves did not have any signs of perforation formation (Fig. 6B,). During the late window (LW) stage in control plants, holes had formed within the perforation sites and were continuously getting larger as more cells within the perforation sites were undergoing PCD and disintegrating (Fig. 6A). The late window stage leaves in AVG treated plants, lacked holes and or any signs of PCD (Fig. 6B). Mature (M) stage leaves were fully perforated within control plants (Fig. 6A), while in AVG treated plants they had few or no perforations (Fig. 6B). Leaves that were produced before treatment (Figs. 6A and B, leaf M1) looked similar and had approximately the same number of perforations.

**AmVPE1 transcript levels in AVG treated and control plants**

Amongst control plants, preperforation and early window stage leaves had the highest *AmVPE1* transcripts compared to all other stages of leaf development. (Fig. 7A). The *AmVPE1* transcript levels were at their lowest in late window and mature stages. Amongst AVG treated plants, *AmVPE1* transcript levels were not significantly different in early preperforation, preperforation, early window and window stage leaves. However, they declined significantly during the late window and mature stages.

The *AmVPE1* transcript levels were also compared between AVG and control plants throughout the 6 stages of leaf development. During preperforation and early window stages, control plants had 2 to 3-fold higher *AmVPE1* transcript levels than AVG treated plants. There was no significant difference in *AmVPE1* transcript levels between AVG treated and control plants in early preperforation, window, late window and mature stages.
AmVPE2 transcript levels in AVG treated and control plants

Amongst control plants, AmVPE2 transcript levels were 1.3 to 2-fold higher during the late window stage than all the other stages of leaf development (Fig. 7B). There were no significant differences in transcript levels between early preperforation, preperforation, early window, window and mature stages. Amongst AVG treated plants, there were no significant differences in AmVPE2 transcript levels throughout leaf development.

There were no significant differences in AmVPE2 transcripts levels between AVG and control plants in all stages of leaf development except the late window stage, where control plants had 1.43-fold higher AmVPE2 transcript levels.

AmActin transcript levels in AVG treated and control plants

AmActin transcripts were constitutively expressed amongst both AVG and control plants (Fig. 7C. Also, in preperforation, window and late window stages, there were no significant differences in AmActin transcript levels between AVG treated and control plants. However, AmActin transcript levels were around 1.23-fold lower in AVG treated plants during early preperforation, early window and mature stages compared to the same developmental stages in control plants.

Discussion

Lace plant VPE transcript levels during leaf development

Percentage identity, phylogenetic and domain analysis suggest that the two lace plant VPEs isolated are vegetative type VPEs. The lace plant VPEs are also more evolutionarily similar to other monocot vegetative type VPEs than dicot vegetative VPEs. Transcript levels suggest that both lace plant VPEs may be involved in PCD during perforation formation in the lace plant. AmVPE1 appears to be involved in the early stages of PCD, before any obvious visible signs of cell degradation (in preperforation stage leaves). AmVPE2 appears to be involved during the later stages of PCD (in window and late window stage leaves), where cellular degradation signs are obvious. Its higher transcript levels during the stages where PCD is actively occurring suggest that AmVPE2 may be involved in the later execution stages of PCD during perforation formation. The higher AmVPE1 transcript levels in preperforation stage leaves and higher AmVPE2 in
transcript levels in the window stage are consistent with the high caspase-1 activity levels reported during these stages of leaf development (Lord et al. 2011). Lord et al. (2011) showed that there is higher caspase-1 activity in preperforation and window stage leaves compared to mature stage leaves. VPEs are responsible for caspase-1 like (YVADase) activity in plants (Hatsugai et al. 2004; Rojo et al. 2004; Misas-Villamil et al. 2013). Taken together, the findings suggest that the observed transcriptional increases of VPEs in the preperforation and window stage leaves may also translate to increased proteolytic activity within these leaves.

In other plant species, VPEs are sometimes known to display increases in transcript levels prior to visible signs of PCD (Iakimova et al. 2013). In apple leaves inoculated with Erwinia amylovora, increases in VPE transcript levels are detected prior to any visible lesion-associated PCD and also after obvious signs of lesion formation (Iakimova et al. 2013). The early-expressed VPEs are thought to be involved in early invisible signs of PCD, signaling/regulation or execution of processes leading to obvious signs of PCD. They may also activate proteases, hydrolases, enzymes responsible for cell degradation and death. VPEs are thought to mediate the maturation of hydrolytic enzymes within the vacuole, which degrade the tonoplast and initiate a proteolytic executing-PCD cascade (Guicciardi et al. 2004; reviewed in Hatsugai et al. 2015). Tonoplast rupture is one of the hallmarks of PCD in the lace plant, and it marks the beginning of rapid cell deterioration (Wright et al. 2009).

Developmentally regulated PCD also occurs as part of leaf senescence. Therefore, it was expected that both AmVPE1 and AmVPE2 transcript levels would be significantly higher during the senescence stage compared to stages in which PCD is not occurring. However, AmVPE1 transcript levels were at their lowest of all leaf developmental stages while AmVPE2 transcripts were not significantly higher than the other stages where PCD is not occurring (Fig. 3). Based on these expression patterns, AmVPE1 and AmVPE2 may not play a role in leaf senescence. However, leaf senescence happens over an extended time period and PCD happens at the end of senescence. The leaves selected were at a specific stage of senescence and both VPEs may be involved at other stages of leaf senescence. An increase in AmVPE1 transcript levels may have happened prior to the PCD signs, as
observed during perforation formation and AmVPE2 could be involved during the later stages where PCD is more rampant. VPEs in other plant species play a role in tissue senescence (Kinoshita et al. 1999; De Michele et al. 2009; Hoeberichts et al. 2007). In *Lilium longiflorum*, VPEs do not seem to be involved in the early stages of flower senescence, but are involved in regulation during the final stages of flower senescence (Battelli et al. 2011). To determine whether both AmVPE1 and AmVPE2 play a role in lace plant leaf senescence, we have to track changes in their transcript levels throughout the process of leaf senescence, instead of a specific senescence stage.

*AmVPE1* and *AmVPE2* transcript levels were significantly higher in the dying cells (PCD cells) than the non-dying cells (NPCD cells; Fig. 4). Suggesting that they may both play a role during the execution of PCD. Generally, *AmVPE2* appeared to have higher transcript levels in leaves than *AmVPE1*. Therefore, it seems to be the more dominant VPE in lace plant leaf tissue, and its transcript expression pattern suggests it plays a more significant role during execution of perforation formation-associated PCD in the lace plant

**VPE activity analysis through JOPD1**

Although the transcript level evidence is compelling, due to the potential post-translational regulation of VPEs, we investigated the activity levels of the VPEs within the two cell types. VPE activity during lace plant PCD was detected using a VPE-activity based probe JOPD1. VPE activity is determined by fluorescent labeling, because it implies the availability and reactivity of VPE active sites. This activity is observed in the vacuoles where VPEs are thought to be active, especially in the PCD types that involve vacuolar rapture, such as the lace plant PCD. Lu et al. (2015) demonstrated that JOPD1 labels all types of VPEs in *Arabidopsis*. The higher VPE activity and fluorescence in PCD cells compared to NPCD cells (Fig. 5) provides more evidence for the involvement of VPEs and VPE activity in lace plant PCD during perforation formation, and more support for the possible central role of VPEs in mediating PCD involving tonoplast rupture. VPE activity probes have been used in leaf extracts and cell cultures before (Misas-Villami et al. 2013; Lu et al. 2015), but to our knowledge, this study is the first to demonstrate the use of VPE activity-based probes in intact tissue sections.
Effect of AVG treatment on AmVPE1 transcript expression

It is thought that ethylene is upstream (within the regulation phase) of VPEs within the plant PCD cascade (reviewed in Rantong and Gunawardena 2015). Lace plant produces high amounts of ethylene during the window stage and inhibition of ethylene biosynthesis stops PCD and perforation formation (Dauphinee et al. 2012). Through our AVG treatment experiments, we showed that reduced ethylene production also correlates with reduced VPE transcript levels. In control plants, the increase in AmVPE1 levels occurs at the beginning of perforation formation and PCD (Figs. 3A and 6). Treatment of plants with AVG seems to prevent the increase in AmVPE1 transcript levels during the preperforation and early window stage (Fig. 7A). Therefore, inhibiting ethylene biosynthesis correlates with hindrance of the increases in AmVPE1 transcript levels during developmental stages where PCD and perforation formation usually occur. The lack of VPE increase results in lack of PCD and perforations. This increase in ethylene levels may be required to trigger increased VPE transcription and execution of PCD in a similar manner proposed by Iakimova et al. (2013).

Effect of AVG treatment on AmVPE2 transcript expression

In control plants, AmVPE2 transcript levels increase significantly during the late window stage (Figs. 3B and 7B). Late window is the stage of leaf development where PCD is vigorously occurring and cells are disintegrating to give rise to holes at perforation sites. This increase is not evident in AVG treated plants. Therefore, the limited ethylene production due to AVG treatment coincides with the lack of increase in AmVPE2 transcript level. Ethylene may also be involved in the signalling leading to the increase in transcriptional upregulation of AmVPE2. The interconnection between ethylene and VPEs is not only unique to the lace plant. Kinoshita et al. (1999) showed that during ripening, treating citrus fruits with ethylene increases VPE transcript levels. Treatment with ethylene also results in increases in VPE transcript levels during plant defense against microbes (Liu et al. 2005).

AmActin transcript levels in AVG treated and control plants

AmActin was constitutionally expressed in both AVG treated and control plants (Fig. 7C). However, during early preperforation, early window and mature stages control plants had
higher transcript levels than AVG treated plants. Ethylene is involved in many other cellular processes unrelated to PCD and its limited production may be affecting other metabolic processes unrelated to PCD including the transcription of housekeeping genes like *AmActin*. However, the 5 µmol/L AVG did not result in any developmental defects within the leaves (except for lack of perforations) and didn’t induce differential expression of *AmActin* transcripts among leaf developmental stages.

Natural substrates cleaved by each of the lace plant VPEs need to be determined and their role during PCD needs to be investigated. The natural substrates would provide insight into whether *AmVPE1* is also involved during the signalling or regulation phase of PCD, since its transcript levels increase early. The ability to transform the lace plant would also be important in order to study the effects of changes in VPE expression through knockout or over-expression lines. The data presented in this study suggest that VPEs are associated with cellular degradation in lace plant PCD and also that ethylene is upstream of VPEs in the plant PCD cascade. Ethylene production also seems to affect the rate and patterns of VPE transcription during lace plant leaf development via PCD. Other components, such as transcription factors, which may interact with VPEs and ethylene within the PCD cascade need to be studied to provide much needed insight into how components within the cascade work together to orchestrate PCD.

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**Author contribution**
Gaolathe Rantong carried out all of the experiments, wrote the first manuscript and made final manuscript revisions. Arunika Gunawardena conceived the study, participated in its design and coordination, and helped in manuscript revisions as well as supervised all experimental work.

Figure legends

Fig. 1. The lace plant.

The lace plant (A) is an aquatic monocot that forms perforations on its leaves through developmentally regulated PCD. It can be grown in sterile conditions in Magenta boxes (B). Early in development, lace plant leaves do not show any signs of perforation formation (C). Eventually cells at the center of a perforation site (PCD cells) undergo PCD, while cells 4-5 layers from the vascular tissue (NPCD cells) remain alive (D). Once the PCD cells have disintegrated, actual holes are formed or perforations result (E). Scale bars (A) = 1.25 cm, B = 1 cm, (C –E) = 100 µm, D = 125 µm and E = 150 µm.

Fig. 2. Alignment and phylogenetic analysis of lace plant VPE amino acid fragments. A. Domains and essential amino acids such as the mature protease domain (blue), essential amino acids (Cys76, His165 and Cys215; red), N-glycosylation site (spring green) and the C-terminal propeptide (orange) were identified. Dots represent amino acid identity and dashes indicate gaps. At the end of each sequence, its percentage identity and similarity with either \textit{Am}VPE1 or \textit{Am}VPE2 are indicated. Accession numbers: AtVPE-gamma (BAA018924.1), AtVPE-alpha (BAA09614.2), AtVPE-beta (BAA09615.1) and AtVPE-delta (BAC65233.1). B. During phylogenetic analysis, the vegetative VPEs from the different species (including lace plant) formed their own clade (highlighted in orange) away from the seed type VPEs (blue). Bar represents the gap separation distance, and the bootstrap values (from a thousand replicates) are indicated above or below each node. Rice asparagine-specific endopeptidase (NP_910213) was used as an outgroup in the Neighbor-Joining tree. Accession numbers of each amino acid sequence are provided in parentheses.
Fig. 3. Changes in *AmVPE1* (A), *AmVPE2* (B) and *AmActin* (C) transcript levels at different stages of leaf development. For each VPE gene at each developmental stage, transcript levels are normalized to the level of *AmActin*, and represent the mean ± SE of ≥12 independent samples. (A) *AmVPE1* transcript levels were higher in preperforation stage leaves compared to all the other stages of leaf development. (B) *AmVPE2* transcripts were higher in window and late window stages, compared to the other stages. (C) *AmActin* transcripts were constitutively expressed throughout leaf development. Similar statistical letters indicate no significant difference at $P > 0.05$. Abbreviations include: early preperforation (EPP), preperforation (PP), early window (EW), window (W), late window (LW), mature (M) and senescence (S).

Fig. 4. Mean Normalized *AmVPE1* and *AmVPE2* Transcript Levels in Dying (PCD) Versus Cells not Undergoing PCD (NPCD Cells).
The PCD and NPCD cells were separated from early window and window stage leaves using a Zeiss PALM Laser Capture Microdissection and Imaging System. Both *AmVPE1* and *AmVPE2* had higher transcript levels in PCD than NPCD cells. *AmActin* transcript levels were not significantly different between PCD and NPCD cells. Means with the same letters are not significantly different ($P > 0.05$). Bars represent SE ($n ≥ 8$).

Fig. 5. VPE activity analysis in PCD and NPCD cells.
VPE activity was detected through an activity-based probe AMS-101. No probe-associated fluorescence was observed in the control (D). In window (B, E and H) and late window (C, F and I) stage leaves, more fluorescence and VPE activity was detected in PCD cells than in NPCD cells (E and F). Dead cells at the centre of the perforation sites did not display any VPE activity. The experiment was repeated three times using 4 replicates in each experiment. A-C = Bright field, D-F = Fluorescence and G-I= Overlay. Scale bars = 70 µm.

Fig. 6. Morphology of AVG treated lace plant compared to control plants.
A leaf layout representing the different developmental stage leaves from control and AVG treated plant (A and B, respectively) was assembled to highlight the differences in perforation numbers between control and AVG treated plants. Control plants had the normal morphology with perforated leaves, while in AVG treated plants most of the leaves produced after treatment lacked perforations. The experiment was repeated four times using ten replicates in each treatment. The leaf developmental stages were early preperforation (EPP, preperforation (PP, early window (EW), window (W, late window (LW) and mature (M). The M1 leaves developed before treatment. Bars = 1.8 cm

Fig. 7. Quantification of AmVPE1, AmVPE1 and AmActin transcripts through QPCR.
Mean normalized AmVPE transcript levels were determined in leaves at different stages of leaf development, in both control and AVG treated plants. During the preperforation and early window stages, AVG treated plants had lower transcript levels than control plants. During the late window stage, control plants had significantly higher AmVPE2 transcript levels than AVG treated plants. For each VPE gene, transcript levels are normalized to the level of AmActin at that developmental stage, and represent the mean ± SE of ≥12 independent samples. AmActin was constitutively expressed throughout leaf development in control and AVG treated plants. However, control plants had higher AmActin transcript levels during early preperforation, early window and mature stages than AVG treated plants. Asterisks highlight developmental stages displaying significant differences (P < 0.05) between treatments. Early preperforation, PP = preperforation, EW = early window, W = window, LW = late window and M = mature

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