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ARTICLE Transcriptional regulation of ethylene and jasmonate mediated defense response in apple (*Malus domestica*) root during *Pythium ultimum* infection

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Apple replant disease (ARD) is a significant economic restraint to the successful re-establishment of new apple orchards on sites previously planted to the same crop. *Pythium ultimum*, an oomycete, is a significant component of the ARD pathogen complex. Although ethylene (ET)- and jasmonic acid (JA)-mediated defense responses are intensively studied in the foliar pathosystem, the transferability of this knowledge to the interaction between a perennial root system and soilborne pathogens is unknown. The aim of this study was to test the hypothesis that the ET/JA-mediated defense response is conserved in roots of tree crops in response to infection by *P. ultimum*. Apple genes with the annotated function of ET/JA biosynthesis, *MdERF* (ethylene response factor) for signaling transduction and a gene encoding a *pathogenesis-related* (*PR*) protein (β -chitinase, the target of ERF) were identified from the apple genome sequences. The transcriptional profiles of these genes during *P. ultimum* infection and after exogenous ET and/or JA treatment were characterized using qRT-PCR. Several genes showed a 10- to 60-fold upregulation in apple root tissue 24-48 h post inoculation (hpi). Exogenous ET and JA treatment exhibited either a positive or negative influence on expression of ET or JA biosynthesis genes, depending upon gene isoforms and the tissue types, while the expression of *MdERF* and the PR protein encoding gene was upregulated by both ET and JA treatment. Our data are consistent with the hypothesis that ET/JA-mediated defense pathways are functional in the root system of perennial tree species defending soilborne pathogens.

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INTRODUCTION

Apple replant disease (ARD) occurs when young trees are planted on a site that has a previous history of apple or closely related species cultivation. Symptoms of the disease can range from a mild uneven growth of young trees to serious growth inhibition or death of young trees. In the absence of control, the effects of ARD are experienced over the entire lifetime of the orchard in the form of decreased fruit yields. The ARD pathogen complex consists of oomycetes including Pythium and Phytophthora and fungi such as Ilyonectria (previously Cylindrocarpon) and Rhizoctonia, and at times acting in concert with the lesion nematode Pratylenchus penetrans.^{1,2} Control of ARD has traditionally relied on pre-plant fumigation of orchard soils; however, future availability of these fumigants could be increasingly restrictive due to environmental concerns. Utilization of rootstock resistance as an important component in managing ARD has not been carefully examined to date, partly due to the complex disease etiology and the difficulty in phenotyping the resistance response.³

Molecular defense responses of plant roots to soilborne necrotrophic pathogens are mostly unknown, and very few studies using root as a model to examine plant–necrotroph interactions.^{4,5} Using the model plant systems and foliar pathogens, ethylene (ET)- and jasmonic acid (JA)-mediated signaling pathways have been demonstrated to be a major part of plant defense reactions against necrotrophic pathogens.⁶⁻¹² However, the transferability of this knowledge from foliar to root tissue is unknown.⁸ The relationship between ET and JA can be antagonistic or synergistic based on different stress signals including pathogen infection.⁷ This complex cross-talk between the two hormonal signaling pathways has been shown to be essential to elicit the appropriate plant reaction in response to pathogen invasion.^{13–15} Both ET- and JA-mediated plant defense responses are merged by activating the expression of *ERF (ethylene response factor)* and *PR (pathogenesis-related)* genes.^{15,16} Recent studies from *Arabidopsis* showed that ERF1, originally thought to have a role only in ethylene signaling, actually responds to both ethylene and JA signals and integrates both hormonal signals to respond to necrotrophic fungal pathogens.^{6,15,17,18} Activated by both hormones, ERF1 regulates plant defense-related genes that have a GCC box on their promoter. Those defense genes appear to function primarily in response to challenges by fungal pathogens and herbivores.^{15,19}

To test the hypothesis that ET/JA-mediated defense response is conserved in both leaf and root tissue, particularly in the root system of perennial tree crops, the transcriptional profiles of genes within the biosynthesis and signaling pathways of ET and JA were studied using apple root tissues infected by *Pythium ultimum*, a component in the ARD pathogen complex. The candidate genes from several gene families were identified from the apple genome sequences.²⁰ These genes included biosynthesis genes for ET (*MdACS: 1-aminocyclopropane-1-carboxylic acid synthase*) and JA (*MdAOS: allene oxide synthase*), ET/JA signal response transcription factor genes (*MdERFs*) and a pathogenesis-related (PR) gene (*MdCHIB*: β -chitinase, the target of ERF). The transcription regulation of these genes to exogenously applied ET and JA was also investigated.

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MATERIALS AND METHODS

Preparation of apple rootstock seedlings and *P. ultimum* inoculum Seeds derived from a Geneva[®] 41 (G.41)²¹ and Malling26 (M.26)²² cross were surface sterilized, germinated and planted in pasteurized SunshineTM potting mix soil (SUN GRO Horticulture Ltd, Bellevue, WA, USA). Seedlings were grown at 23 °C, 95% humidity, under 12/12 h light/dark conditions for 3 weeks.

The *P. ultimum* isolate used in this study was originally isolated from the roots of 'Gala'/M26 apple grown at Moxee, WA, USA.¹ Inoculum of *P. ultimum* was prepared by cultivating in potato carrot broth (20 g of carrots and 20 g of potatoes in 1 L of medium) with two drops of wheat germ oil added per liter of medium. The *P. ultimum* cultures were grown in the broth in Petri dishes at 22 °C for 2–3 weeks. Oospores and mycelium were separated from the media by passing through a double layer of cheese cloth and the resultant mat was ground in a blender for 30 s. Spores and hyphal fragments were resuspended in 2% methyl cellulose to give a final concentration of approximately 2000 colony forming units mL^{-1} .

The inoculation of seedlings with *P. ultimum* was performed by dipping 70%–80% of the entire root system into the inoculum for 5 s; then inoculated seedlings were planted into pasteurized SunshineTM potting mix. The inoculated seedlings were grown in an environmental growth chamber at 23 °C, 95% humidity, under a 12/12 h light/dark photoperiod. Plants root tissues were collected at 0, 2, 4, 8, 24, 48, 72 and 96 h post inoculation (hpi) and collected tissues were frozen in liquid nitrogen. Root tissue of twenty seedlings was collected and pooled for each time point per treatment. The experiment was repeated twice. Frozen root tissues were dipped into 2% methyl cellulose solution and planted into pasteurized SunshineTM potting mix.

ET and JA treatment

Seedlings were prepared as described above. Ethylene treatment was performed by purging 110 ppm ET (background with air) into a 3.8 L-size jar for 30 s. JA treatment was performed by pouring 500 μ M methyl jasmonate (Sigma-Aldrich, St. Louis, MO, USA) in 0.177% Tween 20 into 2.25 inch square pots (Jiffy Products Ltd, Shippagan, Canada) until the pasteurized SunshineTM potting mix soil was saturated. The seedlings were treated at room temperature for 6 h at which time plants were extracted from the potting soil. Root and shoot tissue of 15–20 seedlings were collected and pooled per treatment. The experiment was repeated twice. Roots were excised from shoots and separately placed in liquid nitrogen. Frozen tissues were stored at -80 °C.

Primer design and phylogenetic tree construction

Forward and reverse primers for *MdACSs*, *MdAOSs*, *MdERFs*, *MdCHIB* and *MdActin* genes (Table 1) were designed using web-based Primer3plus software (http://www.bioinformatics.nl/cgi-bin/primer3plus/primer3plus.cgi) and an IDT oligo analyzer (http://www.idtdna.com/analyzer/Applications/

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OligoAnalyzer/). Where possible, an optimum annealing temperature of 60 °C, a GC content of 40%–60%, an amplicon length of 150–180 bp and a primer length of 20 bp were applied. Multisequence alignment software ClustalW (http://www.ebi.ac.uk/clustalw) and BLAST (http://www.ncbi.nlm .nih.gov/BLAST/) were used to choose divergent regions at or close to 3' untranslated region for gene-specific primer design. Amplicons were sequenced to confirm identity of the specific amplified genes. Clustal W2 from EMBL-EBI (http://www.ebi.ac.uk/embl/) website and the Jalview 2.6.1 program were used to identify similarities between amino-acid sequences and to construct a phylogenetic tree. To calculate the distance between sequences, BLOSUM62 was used. Neighbor joining method was used for the phylogenetic tree construction.

RNA isolation and real-time gRT-PCR

The frozen tissue samples were ground to a fine powder in liquid nitrogen and total RNA was isolated following the modified method described by Gasic et al.²³ The total RNA was quantified using a ND 1000 Nanodrop spectrophotometer (NanoDrop Technologies, Wilmington, DE, USA) and the RNA quality was verified by agarose gel electrophoresis. Total RNA was treated with DNase I (Qiagen, Valencia, CA, USA) and then purified with RNeasy cleanup columns (Qiagen, Valencia, CA USA). Two micrograms of DNase-treated RNA was used to synthesize first-strand cDNA using SuperScriptTM II reverse transcriptase (Invitrogen, Grand Island, NY, USA) and poly dT (Operon, Huntsville, AL, USA) as the primer. The cDNA was diluted 20 times and 0.6 µL aliquot was used in a 15 µL quantitative PCR (qPCR) reaction mix: 0.45 µL SYBR Green I dye (Invitrogen, Grand Island, NY, USA), 1X iTaq buffer (Bio-Rad, Hercules, CA, USA), 0.2 mM dNTP (Applied Biosystem, Foster City, CA, USA), 2.5 mM MgCl₂, 0.3 units of iTaq DNA polymerase (Bio-Rad, Hercules, CA, USA) and 0.2 μ M forward/ reverse primer (IDT). Real-time qPCR amplification and detection was conducted using an iQ5 real-time qPCR detection system (Bio-Rad Lab, Hercules, CA) and the following protocol: cycle conditions of 3 min at 95 °C, 40 cycles of 10 s at 95 °C and 30 s at 59 °C. Dissociation curves were run for MdACSs, MdAOSs, MdERFs, MdCHIB and MdActin primers to determine the presence of any non-specific amplification. The relative gene expression was measured using the 0 h root tissue (for P. ultimum infection) or no-hormone treated control (for ET and JA transcriptional regulation) as the calibrator. 'No reverse transcriptase' and 'no template' negative controls were included in PCR amplification. Two sets of root samples (biological replicates) were used to obtain two independent total RNA sample sets, and subsequently converted into two separate cDNA sample sets. For each cDNA sample set, PCR reactions were performed twice. Therefore, four values of PCR amplifications (four data sets per sample: two biological replicate, each with two technical replicates) were performed from two independent biological samples. PCR amplification was carried out in triplicate in a 96-well plate. The target gene expression was normalized to that of the internal reference gene (*MdActin*) using the $2^{-\Delta\Delta Ct}$ method (the comparative C_t method).^{24,25}

Gene name	Gene model from apple genome	Chromosomal location	Primer sequences
MdACS	MDP0000130748	Chr 1: 9.86 Mb	AAT CGT TAA TGA AGT GAA GCT C
			GAC TTT TCA CTT GTG CTT GT
MdACS	MDP0000262827	Chr 1: 9.88 Mb	CAA GAA CAA GCA CAA GTG AA
			GGT CCC CAA TAC TCT TTA AGT
MdACS	MDP0000435100	Chr 7: 14.19 Mb	GCA ATT GTG CCG AGG A
			GGG AGA TCA GTC ACT TCT CTA
MdAOS	MDP0000132456	Chr 2: 27.53 Mb	GAC TAC TTC TAC AAC CAA GGC
			GGT GGT GTC AAA GAG AAT GG
MdERF1	MDP0000127134	Chr 13: 25.73 Mb	CAG TTG AAA GAG TCC GCA A
			CAA ACA CCA CCA CAT TCT CTA
MdERF1	MDP0000235313	Chr 16: 18.14 Mb	GTA GAG AGA GTC CGC GAA T
			CAA ACA CCA CCA CAT TCT CTA
MdERF	MDP0000880063	NA	AGC TGC TTT TTC CAT GAG G
			GAT TCC GTC CTG GTC ACA
MdCHIB	MDP0000430546	Chr 4: 1.09 Mb	GTC ACT CAG GCA TTC TTC G
			CAT GGG TGA CAT GAG CAA A
MdActin	MDP0000752428	Chr 4: 12.33 Mb	GTC GTA CTA CTG GTA TCG TT
			TCA TAG TCA AGA GCA ATG TA

RESULTS

The apple root symptom from P. ultimum infection

A representative image of *P. ultimum* infected apple root tissues is shown in Figure 1. Compared with un-inoculated controls showing a white and healthy root tissue (Figure 1a), tissue necrosis with dark brown coloration throughout the root was typically observed on *P. ultimum* infected root (Figure 1b). It was also shown that the actively growing area of roots, especially the tips of emerging lateral roots or primary roots, are highly susceptible to infection by *P. ultimum* (Figure 1b). The symptoms represented in this figure were routinely observed.

Expression profiles of ET and JA biosynthesis genes in response to *P*. *ultimum* infection

To identify the members of the gene family for ET and JA biosynthesis that are induced in response to the P. ultimum infection, expression patterns of apple MdACS and MdAOS genes were examined. The phylogenetic tree for these fifteen MdACS genes was constructed by comparing their encoded peptide sequences and constructing a phylogenetic tree (Figure 2). The phylogenic analysis divided MdACSs into three subgroups: Groups I, II and III (Figure 2). Among the 15 MdACS isoforms, three genes (MDP0000130748, MDP0000262827 and MDP0000435100) exhibited a 6-60 time increase of transcript abundance within 48 hpi of P. ultimum (Figure 3a–3c). The expression data of the MdACSs that were not affected by *P. ultimum* infection are provided in Supplementary Fig. S1. Among the three P. ultimum infection induced MdACS genes, MDP0000435100 showed the highest response at a time point 24 h after inoculation (Figure 3c). All three MdACS genes that responded to P. ultimum belong to the same sub-group (group I), while the well-studied fruit ripening related ethylene biosynthesis genes



Figure 1. The observed symptoms of *P. ultimum* infected apple root tissue. (**a**) Un-inoculated control shows a healthy root tissue with light color. (**b**) *P. ultimum*-infected root show a dark brown necrosis. The close-up images on tips of emerging lateral root in un-infected control (**a**) and *P. ultimum*-infected root system (**b**) also demonstrate a difference in coloration between healthy tissue (**a**) and necrosis tissue (**b**).

(*MdACS3* and *MdACS1*) belong to group II and III, respectively (Figure 2) and the expression of these fruit ripening-related genes was not affected by *P. ultimum* (Supplementary Fig. S1). Subgroups of the *MdACS* family and their differential gene expression in ripening and *P. ultimum* inoculation indicated that MdACS isoforms specifically participate in distinct plant processes.

Similarly, the expression profiles of members from the *MdAOS* gene family (*MDP0000132456*, *MDP0000150140*, *MDP0000424398*, *MDP0000198152* and *MDP0000225501*) for JA biosynthesis in apple root tissue were investigated in response to inoculation with *P. ultimum*. Among the five genes examined, one *MdAOS* (*MDP0000132456*) showed strong upregulation in root tissues inoculated with *P. ultimum* (Figure 3d). Peak expression of this gene was observed 48 hpi with at least a 40-fold increase in transcript abundance, which corresponded with the expression pattern of the three *MdACS* genes (Figure 3).

Identification of P. ultimum responsive MdERFs and MdCHIB

Four ERF genes (*MDP0000235313*, *MDP0000167207*, *MDP0000127134* and *MDP0000880063*) in apple showed an upregulation upon inoculation with *P. ultimum* (Figure 4a–4c). Based on genomic location and DNA/peptide sequence identity (data not shown), two of the *MdERF1* genes (*MDP0000167207* and *MDP0000127134*) were considered to be the same gene, and therefore only *MDP0000127134* was included in further studies. Timing of the observed upregulation in the expression of the *MdERF* genes corresponded with the ET/JA hormone biosynthesis gene expression and was highly induced 24–48 hpi (Figures 3 and 4). Additional *MdERFs* that were not upregulated are described in Supplementary Fig. S3.

Bioinformatic analysis demonstrated that the upregulated apple MdERFs share a substantial similarity in peptide sequences with their homologous *Arabidopsis* AtERFs: MdERF1 (MDP0000235313 and MDP0000127134) possessed 55%–56% identity and 62% similarity to AtERF1 (At3g23240);¹⁹ MdERF (MDP0000880862) had 43% identity and 58% similarity to AtERF (At1g04370)²⁶ (Figure 5a). These three MdERFs and two AtERFs showed a consensus AP2/ERF (APETALA2/ET response factor) domain²⁷ (Figure 5a). The phylogenetic tree indicated that MdERF1 (MDP0000235313 and MDP0000127134) is closely related to the well-characterized pathogen-responsive AtERF1 (At3g23240)¹⁹ and ORA59 (At1g06160)²⁸ (Figure 5b).

An apple chitinase gene (*MdCHIB*: *MDP0000430546*) was also identified to be upregulated in response to infection by *P. ultimum*. *CHIB* (*CHITINASEB*) has been known to be a *pathogenesis-related* (*PR*) gene and its promoter region is a target of the ERF transcription factor.^{16,19,28} *MdCHIB* (*MDP0000430546*) exhibited a strong upregulation in response to *P. ultimum* infection, particularly in the period from 48 to 72 hpi (Figure 4d).

Transcriptional regulation of apple *P. ultimum*-activated genes to exogenous ET and JA treatment

To gain more insight into the ET/JA hormonal regulation of *P. ultimum* infection-activated genes, the expression patterns of *MdACS*, *MdAOS*, *MdERF* and *MdCHIB* in response to exogenous ET/JA treatment were studied. All three *MdERF* genes were upregulated by either ET or JA (Figure 6a–6f). Moreover, when ET and JA were applied in combination, a synergistic response at the gene expression for all *MdERF* genes (i.e. a higher fold changes) was observed (Figure 6a–6f). Although regulation of these two *MdERF1* was similar, the expression of *MDP0000235313* exhibited greater sensitivity to ethylene than did *MDP0000127134* (Figure 6a–6d). Unlike the other two *MdERFs*, *MDP0000880063* was induced to a higher level by JA than by ethylene (Figure 6e–6f). ET and JA appeared to act synergistically in the induction of the *MdCHIB* gene expression (Figure 6q–6h).

The expression of the three *MdACS* genes was regulated by ET and JA signals both antagonistically and synergistically. For two of

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Figure 2. Phylogenetic tree of all apple 15 MdACS genes. Based on deduced amino-acid sequence similarity, MdACSs were divided into three groups: group I, II, and III. Three MdACSs that were upregulated by the *P. ultimum* infection are in bold. Md, *Malus domestica*.



Figure 3. Gene expression profile of three apple MdACS and MdAOS in response to P. ultimum in apple root: (a) MdACS (MDP0000130748); (b) MdACS (MDP0000262827); (c) MdACS (MDP0000435100); (d) MdAOS (MDP0000132456). Each data point represents the means with standard error.

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Figure 4. Gene expression profiles of three apple *MdERFs* and *MdCHIB* in response to *P. ultimum* in apple root: (a) *MdERF1* (*MDP0000235313*); (b) *MdERF1* (*MDP0000127134*); (c) *MdERF1* (*MDP0000880063*); (d) *MdCHIB* (*MDP0000430546*). Each data point represents the means with standard error.

the *MdACS* genes (*MDP0000130748* and *MDP0000262827*), ET acted as a negative regulator while JA acted as a positive regulator of their expression (Figure 7a–7d). This antagonistic regulation between the two hormones was observed in both root and shoot tissues (Figure 7a–7d). The expression of the third *MdACS* (*MDP0000435100*) was upregulated individually by ET or JA, and two hormones acted synergistically, also its expression was much higher in root than in shoot tissues (Figure 7e–7f). In response to ET and JA treatments, the expression of *MdAOS* (*MDP0000132456*) showed significantly greater upregulation in roots (approximately 70-fold increase at transcript abundance) than in shoots, but a synergistic response was only observed in shoot tissue (Figure 7g–7h). These results added evidence that the expression of *P. ultimum* response genes is directly or indirectly regulated by both ET and JA hormonal signals.

DISCUSSION

The role of ET/JA mediated defense response in plant root, especially in perennial tree species, is still mostly unknown,^{29,30} though ET/JA signaling pathways in a plant defense response in leaf tissues against necrotrophic pathogens has been well established, mostly from foliar disease systems of model plants such as Arabidopsis and tobacco.^{8,31–33} Findings from the current study demonstrate that the ET/JA signaling pathways are functional in apple root in response to the *P. ultimum* infection. Our data showed that infection by *P. ultimum* activated ET/JA signaling-related genes, including *MdACS* (ethylene biosynthesis gene), *MdAOS* (JA biosynthesis gene), *MdERF* (a transcription factor gene that responds to both ET/JA hormonal signals) and *MdCHIB* (*CHITINASEB* is a target of ERF) in the apple root system.

Although the transcriptional regulation of genes for ET and JA biosynthesis and signaling is just one layer of their function, results from this study provide the essential information for further elucidation of their regulating roles in defense response. The coordinated regulation patterns of genes in both ET and JA biosynthesis pathways in response to P. ultimum inoculation suggested that both hormones are required to activate an effective defense reaction in root tissues. Our data indicated that most ET/ JA biosynthesis genes, except MdACS (MDP0000435100), exhibited peak induction at 48 hpi. Two out of three MdERFs also showed peak expression at 48 hpi, while its downstream PR protein encoding gene MdCHIB displayed maximum expression at 72 hpi. The coordinately upregulated expression of the genes suggested that a functional ET/JA-mediated defense pathway (from ET/JA biosynthesis and signaling transduction to antimicrobial activity) exists in apple roots.

Cross-talk among hormone signaling networks during plant defense has been well elucidated in which ET and JA signaling pathways regulate and modulate each other to produce effective defense responses.^{8–10,34,35} ET and JA independently activate not just separate sets of genes, they also synergistically activate *PR* genes (including *PDF1.2* and *CHIB*) through ERF in response to necrotrophic pathogens.^{6,8,34,35} In the JA signaling pathway, COI1 (JA receptor) receives the JA signal and initiates degradation of JAZ (JA response repressor) through ubiquitination and proteasome activity.³⁶ The removal of JAZ boosts EIN3/EIL1 activity which causes induction of *ERF* and consequently induces *CHIB*. Ethylene signaling is incorporated through enhancing EIN3/EIL1 stability. Since EIN3/EIL1 protein has a short-life cycle, the ethylene signal is essential for its build-up in the nucleus.^{37,38} Two *MdACSs*

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а	MDP0000127134	MRCHKYTN TXXVFQAENKNTPILQMNYSTFD SPNSNLSPESSFGSSFSWDDGLKFGNNSL	60
	MDP0000235313	MNYSTED SPNSNLSPESSEGSSESWDDGLKEGNNSL	36
	At3g23240	MDPFLIQSPFSGFSPEYSIGSSPDSFSSSSSNNYSL	36
	MDP0000880063		
	At1g04370		
	MDP0000127134	PFNENDSEEMLLYGLISEGTOEITSSVSVFANPIKEEEVSSASEEENPKKEKSVRGVRR	120
	MDP0000235313	PFNENDSEEMLLYGLISEATQEITSSFFVSANPIKEEEVSSACEEENLKKEKSVRGVRRR	96
	At3g23240	PFNENDSEEMFLYGLIEQSTQQTYIDSDSQDLPIKSVSSRKSEKSVRCVRRR	88
	MDP0000880063	QKDVKYRGKDKEEGQKDV	23
	At1g04370	AEQGKSSGSGGGAEQGKYRGWRR	26
	MDP0000127134	PWGKFAAEIRDSTRHGIRVWLGTFDSAEAAALAYDQAAFSMRGSAAVLNFP <mark>VERVRKSLR</mark>	180
	MDP0000235313	PWGKFAAEIRDSTRHGIRVWLGTFDSAEAAALAYDOAAFSMRGSXAILNFP <mark>VERVRESLR</mark>	156
	At3g23240	PWGKFAAEIRDSTRNGIRVWLGTFESAEEAALAYDOAAFSMRGSSAILNFS <mark>AERVO</mark> ESLS	148
	MDP0000880063	PWGKFAAEIRDSTROGARLWLGTFNTAEEAARAYDRAAFSMRGPLAILNFP <mark>IEYDLK</mark>	80
	At1g04370	PWGKYAAEIRDSRKHGERVWLGTFDTAEDAARAYDRAAYSMRGKAAILNFP <mark>HEYNMG</mark>	83
	MDP0000127134	XINYGTSD LEGCSPVVALKRKHSMRRKTGNTKSKVDRDVRIENVVVFEDLGIDYLE	236
	MDP0000235313	EINYGTLD LEGCSPVMALKRKHSMRRKTGFKKSQVDRDVMIENVVVFEDLGTDYLE	212
	At3g23240	EIKYTYEDGCSPVVALKRKHSMRRRMTNKKTKDSDFDHRSVKLDNVVVFEDLGEQYLE	206
	MDP0000880063	TRTESGREIFEFEC LDDSVLE	126
	At1g04370	QQVFEFEY LDDSVLD	112
	MDP0000127134	ELLNSSESTSTNATPNW 253	
	MDP0000235313	ELLNSTENTSTSATPNW 229	
	At3g23240	ELLGSSENSGTW 218	
	MDP0000880063	DLLDF-DNHTKTNEEKQIK 144	
	At1g04370	ELLEYGENYNKTHNINMGKRQ 133	



Figure 5. Sequence similarity shared between MdERFs and AtERFs. (a) Alignment of deduced amino-acid sequence of two Arabidopsis AtERFs (At3g23240 and At1g04370) and three apple MdERFs (MDP0000127134, MDP0000235313 and MDP0000880063). AP2/ERF domain is indicated by reverse lettering. (b) Phylogenetic analysis of MdERFs from this study and AtERFs that were reported as necrotrophic pathogen and JAinduced genes. Three MdERFs that were upregulated by *P. ultimum* infection are in bold.

(MDP0000130748 and MDP0000262827) and MdAOSs (only in root) were shown to be induced by JA and repressed by ET application and an antagonistic regulation by combinational ET/JA treatments. On the other hand, the expression of MdERF and MdCHIB, were upregulated by both individual ET and JA treatments or synergistically by ET/JA treatment in combination. These observations suggest that first, complicated feedback regulation existed at ET/JA

biosynthesis and second, both hormones act synergistically for the signal transduction through MdERF and downstream defense actions in both apple root and shoot tissues.

For the regulation of ET/JA biosynthesis genes, our results based on exogenous ET and JA treatment assays indicated that JA always acts as a positive regulator, while ET acts as either positive or negative feedback signals. The contrasting action of ET in shoots and

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Figure 6. Gene expression profile of two apple *MdERF1s* with ET and JA treatment: (**a** and **b**) *MdERF1* (*MDP0000235313*); (**c** and **d**) *MdERF1* (*MDP0000127134*); (**e** and **f**) *MdERF* (*MDP0000880063*); (**g** and **h**) *MdCHIB* (*MDP0000430546*); (**a**, **c**, **e** and **g**) under-ground tissue (roots); (**b**, **d**, **f** and **h**) above-ground tissue (shoot). ET: 110 ppm ethylene only; JA: 500 μ M methyl jasmonate only; ET/JA: ethylene and methyl jasmonate together. Each data point represents the means with standard error.

roots could be due to other factors such as Ca²⁺ signaling and mitogen-activated protein kinase, which affect ET-mediated transcriptional regulation in addition to the major regulatory mechanism of ubiqutin-mediated protein degradation.^{15,37} For JA, it has been known that JA biosynthesis genes, including LOX (Lipoxygenase), AOS, AOC (Allene Oxide Cyclase), OPR3 (OPDA

reductase 3) and *ACX (AcyI-CoA Oxidase)*, can be activated through the SCF^{COI1}-JAZ-mediated positive feedback loop.^{15,39} Between root and shoot tissues, very similar gene expression patterns were observed for all genes studied except *MdAOS*, suggesting the function of similar molecular components in the defense response in both shoot and root tissues. -

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Figure 7. Gene expression profile of three apple *MdACSs* with ET and JA treatment: (**a** and **b**) *MdACS* (*MDP0000130748*); (**c** and **d**) *MdACS* (*MDP0000262827*); (**e** and **f**) *MdACS* (*MDP0000435100*); (**g** and **h**) *MdAOS* (*MDP0000132456*); (**a**, **c**, **e** and **g**) under-ground tissue (roots); (**b**, **d**, **f** and **h**) above-ground tissue (shoot). ET: 110 ppm ethylene only; JA: 500 μ M methyl jasmonate only; ET/JA: ethylene and methyl jasmonate together. Each data point represents the means with standard error.

A simplified model of ET/JA biosynthesis and signaling in apple roots in response to *P. ultimum* infection was shown in Figure 8 based on the observations from this study. Infection by *P. ultimum* triggers ET/JA biosynthesis by activating *MdACS* and *MdAOS* in the apple root. ET/JA signaling pathways are activated when the ET/JA hormonal receptor perceives signals, which synergistically upregulate the expression of *MdERF1* preassembly through SCF^{COI1}-JAZ proteasome activity and EIN3/EIL1 transcriptional activity. MdERF1, as a transcription factor, activates the expression of *PR* genes such as *MdCHIB* by binding to their promoter



Figure 8. The proposed model of hormonal regulation (by ET and JA) on transcription of *P. ultimum* response genes in apple root. Accumulated ET and JA in response to the *P. ultimum* infection activate the *MdERF1* expression through SCF^{COI1}-JAZ proteasome activity for subsequent activation of *MdCHIB*. ET and JA also can regulate their own biosynthesis pathways as either positive or negative feedback regulation.

region. JA not only activates defense-related genes, but JA is also involved in feedback regulation of JA biosynthesis by acting as a positive regulator in the apple root system. This study serves as a proof of concept that the knowledge of plant defense response to necrotrophs, which were primarily elucidated using foliar pathogens of *Arabidopsis*, seems to be transferable to the interaction between root of perennial crops and soil borne necrotrophic pathogens.

COMPETING INTERESTS

The authors declare no conflicts of interest.

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REFERENCES

- Mazzola M. Elucidation of the microbial complex having a causal role in the development of apple replant disease in Washington. *Phytopathology* 1998; 88: 930–938.
- 2 Tewoldemedhin YT, Mazzola M, Botha WJ, Spies CF, McLeod A. Characterization of fungi (*Fusarium* and *Rhizoctonia*) and oomycetes (*Phytophthora* and *Pythium*) associated with apple orchards in South Africa. *Eur J Plant Pathol* 2011; **130**: 215–229.
- 3 Zhu Y, Fazio G, Mazzola M. Elucidating the molecular responses of apple rootstock resistant to ARD pathogens: challenges and opportunities for development of genomics-assisted breeding tools. *Hort Res* 2014; **1**: 14043.
- 4 Li C, Shao J, Wang Y, Li W, Guo D *et al.* Analysis of banana transcriptome and global gene expression profiles in banana roots in response to infection by race 1 and tropical race 4 of *Fusarium oxysporum* f. sp. cubense. *BMC Genomics* 2013; 14: 851.

- 5 Millet YA, Danna CH, Clay NK, Songnuan W, Simon MD *et al.* Innate immune responses activated in *Arabidopsis* roots by microbe-associated molecular patterns. *Plant Cell* 2010; 22: 973–990.
- 6 Shinshi H. Ethylene-regulated transcription and crosstalk with jasmonic acid. *Plant Sci* 2008; **175**: 18–23.
- 7 Kazan K, Manners JM. Jasmonate signaling: toward an integrated view. *Plant Physiol* 2008; **146**: 1459–1468.
- 8 Okubara P, Paulitz T. Root defense responses to fungal pathogens: a molecular perspective. *Plant Soil* 2005; 274: 215–226.
- 9 Glazebrook J. Contrasting mechanisms of defense against biotrophic and necrotrophic pathogens. *Annu Rev Phytopathol* 2005; **43**: 205–227.
- 10 Pieterse CM, van der Does D, Zamioudis C, Leon-Reyes A, van Wees SC. Hormonal modulation of plant immunity. Annu Rev Cell Dev Biol 2012; 28: 489–521.
- 11 Ballare CL. Jasmonate-induced defenses: a tale of intelligence, collaborators and rascals. *Trends Plant Sci* 2011; **16**: 249–257.
- 12 Robert-Seilaniantz A, Grant M, Jones JD. Hormone crosstalk in plant disease and defense: more than just jasmonate–salicylate antagonism. *Annu Rev Phytopathol* 2011; 49: 317–343.
- 13 Acosta IF, Farmer EE. Jasmonates. The Arabidopsis Book 2010; 8: e0129.
- 14 Mengiste T. Plant immunity to necrotrophs. Annu Rev Phytopathol 2012; 50: 267–294.
- 15 Wasternack C, Hause B. Jasmonates: biosynthesis, perception, signal transduction and action in plant stress response, growth and development. An update to the 2007 review in *Annals of Botany. Ann Bot* 2013; **111**: 1021– 1058.
- 16 Solano R, Stepanova A, Chao Q, Ecker JR. Nuclear events in ethylene signaling: a transcriptional cascade mediated by ETHYLENE-INSENSITIVE3 and ETHYLENE-RESPONSE-FACTOR1. Genes Dev 1998; 12: 3703–3714.
- 17 Berrocal-Lobo M, Molina A, Solano R. Constitutive expression of ETHYLENE-RESPONSE-FACTOR1 in Arabidopsis confers resistance to several necrotrophic fungi. Plant J 2002; 29: 23–32.
- 18 Wasternack C. Jasmonates: an update on biosynthesis, signal transduction and action in plant stress response, growth and development. Ann Bot 2007; 100: 681–697.
- 19 Lorenzo O, Piqueras R, Sánchez-Serrano JJ, Solano R. ETHYLENE RESPONSE FACTOR1 integrates signals from ethylene and jasmonate pathways in plant defense. *Plant Cell* 2003; **15**: 165–178.
- 20 Velasco R, Zharkikh A, Affourtit J, Dhingra A, Cestaro A et al. The genome of the domesticated apple (*Malus x domestica* Borkh.). *Nat Genet* 2010; **42**: 833– 839.
- 21 Fazio G, Aldwinckle HS, Robinson TL, Cummins J. (314) Geneva[®] 41: a new fire blight resistant, dwarf apple rootstock. *HortScience* 2005; **40**: 1027.
- 22 Wertheim SJ. Rootstock Guide: Apple, Pear, Cherry, European Plum. Wilhelminadorp: Fruit Research Station; 1998.
- 23 Gasic K, Hernandez A, Korban S. RNA extraction from different apple tissues rich in polyphenols and polysaccharides for cDNA library construction. *Plant Mol Biol Rep* 2004; 22: 437–438.
- 24 Livak KJ, Schmittgen TD. Analysis of relative gene expression data using real-time quantitative PCR and the $2^{-\Delta\Delta CT}$ method. *Methods* 2001; **25**: 402–408.
- 25 Schmittgen TD, Livak KJ. Analyzing real-time PCR data by the comparative C_T method. *Nat Protoc* 2008; **3**: 1101–1108.
- 26 McGrath KC, Dombrecht B, Manners JM, Schenk PM, Edgar Cl et al. Repressorand activator-type ethylene response factors functioning in jasmonate signaling and disease resistance identified via a genome-wide screen of *Arabidopsis* transcription factor gene expression. *Plant Physiol* 2005; **139**: 949–959.
- 27 Okamuro JK, Caster B, Villarroel R, van Montagu M, Jofuku KD. The AP2 domain of APETALA2 defines a large new family of DNA binding proteins in *Arabidopsis. Proc Natl Acad Sci USA* 1997; **94**: 7076–7081.
- 28 Pre M, Atallah M, Champion A, de Vos M, Pieterse CM et al. The AP2/ERF domain transcription factor ORA59 integrates jasmonic acid and ethylene signals in plant defense. Plant Physiol 2008; 147: 1347–1357.
- 29 Vijayan P, Shockey J, Lévesque CA, Cook RJ, Browse J. A role for jasmonate in pathogen defense of Arabidopsis. Proc Natl Acad Sci USA 1998; 95: 7209– 7214.
- 30 Geraats BP, Bakker PA, van Loon LC. Ethylene insensitivity impairs resistance to soilborne pathogens in tobacco and Arabidopsis thaliana. Mol Plant Microbe Interact 2002; 15: 1078–1085.
- 31 Wang KL, Li H, Ecker JR. Ethylene biosynthesis and signaling networks. Plant Cell 2002; 14(Suppl 1): S131–S151.
- 32 Turner JG, Ellis C, Devoto A. The jasmonate signal pathway. *Plant Cell* 2002; **14**(Suppl 1): S153–S164.
- 33 Broekaert WF, Delauré SL, de Bolle MF, Cammue BP. The role of ethylene in hostpathogen interactions. *Annu Rev Phytopathol* 2006; **44**: 393–416.

- Dong X. SA, JA, ethylene, and disease resistance in plants. *Curr Opin Plant Biol* 1998;
 1: 316–323.
- 35 Feussner I, Wasternack C. The lipoxygenase pathway. Annu Rev Plant Biol 2002; 53: 275–297.
- 36 Wager A, Browse J. Social network: JAZ protein interactions expand our knowledge of jasmonate signaling. *Front Plant Sci* 2012; **3**: 41.
- 37 Zhao Q, Guo HW. Paradigms and paradox in the ethylene signaling pathway and interaction network. *Mol Plant* 2011; 4: 626–634.
- 38 An F, Zhao Q, Ji Y, Li W, Jiang Z et al. Ethylene-induced stabilization of ETHYLENE INSENSITIVE3 and EIN3-LIKE1 is mediated by proteasomal degradation of EIN3 binding F-box 1 and 2 that requires EIN2 in Arabidopsis. Plant Cell 2010; 22: 2384–2401.
- 39 Chung HS, Koo AJ, Gao X, Jayanty S, Thines B *et al*. Regulation and function of Arabidopsis JASMONATE ZIM-domain genes in response to wounding and herbivory. *Plant Physiol* 2008; **146**: 952–964.

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