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31 SUMMARY

Bacterial wilt caused by the bacterial pathogen Ralstonia solanacearum is one of the most devastating crop diseases worldwide. The molecular mechanisms controlling the early stages of *R. solanacearum* colonization in the root remain unknown. In this study we established four stages in the early interaction of the pathogen with Arabidopsis roots and determined the transcriptional profiles of these stages of infection. A total 2698 genes were identified as differentially expressed genes during the initial 96h after infection, with the majority of changes in gene expression occurring after pathogen-triggered root hair development was observed. Further analysis of differentially-expressed genes indicated sequential activation of multiple hormone signaling cascades, including abscissic acid (ABA), auxin, jasmonic acid (JA), and ethylene (ET). Simultaneous impairment of ABA receptor genes increased plant sensitivity to R. solanacearum, but did not affect primary root growth inhibition, root hair and lateral root formation caused by R. solanacearum. This indicates that ABA signaling positively regulates root defense to R. solanacearum. Moreover, transcriptional changes of genes involved in primary root, lateral root and root hair formation exhibited high temporal dynamics upon infection. Taken together, our results suggest that successful infection of R. solanacearum on roots is a highly programmed process involving in hormone crosstalk.

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INTRODUCTION

Ralstonia solanocearum, a soil-borne phytopathogen, causes devastating bacterial wilt disease on crops and leads to huge economical loss (Mansfield et al., 2012). The bacterium enters into the root epidermis through natural openings or wounds, crosses the cortex and endodermis and finally reaches the root xylem. In the xylem, R. solanacearum starts extensive colonization, spreading to the aerial part of the infected plant along the vascular system and finally kills the host by blocking water transport from root to shoot, which causes the wilting symptoms (Genin and Denny, 2012). Due to its wide host range, long persistence in soil and water and broad geographical distribution, R. solanacearum was ranked as the second most important bacterial plant pathogen (Mansfield et al., 2012).

The interaction between R. solanacearum and Arabidopsis has been successfully used for more than twenty years as a model to study plant defense (Deslandes et al., 1998). However, our knowledge about the molecular mechanisms used by Arabidopsis to defend against R. solanacearum is still limited. RRS1-R is the only R. solanacearum resistance gene cloned from Arabidopsis and encodes a Toll-IL-1 receptor-nucleotide binding site-leucine rich repeat (TIR-NB-LRR) resistance protein with a C-terminal WRKY DNA-binding motif (Deslandes et al., 2002). In the absence of PopP2, an effector from R. solanacearum GMI1000, RRS1-R forms heterodimer complex with RPS4, another NB-LRR protein, localizes in the nucleus and binds DNA through WRKY domain. When PopP2 is delivered into the host cell through the Type Three Secretion System (TTSS), it directly interacts with the RPS4/RRS1-R resistance complex and acetylates WRKY domain of RRS1-R through its acetyltransferase activity, blocking RPS4/RRS1-R DNA-binding activity and activating RPS4-mediated plant resistance (Le Roux et al., 2015; Sarris et al., 2015).

It is widely recognized that the phytohormones salicylic acid (SA), jasmonate (JA) and ethylene (ET) play a determinant role in plant defense to diverse

pathogenic insects, bacteria and fungi. However it is still not clear what is the precise role of these hormones in response to R. solanacearum. Arabidopsis mutants deficient in biosynthesis or signaling of SA, JA and ET have been used to investigate their sensitivity to R. solanacearum, which has led sometimes to contradictory results. For instance, while an increase or decrease of endogeneous SA levels did not alter plant sensitivity to R. solancearum (Hirsch et al., 2002), depletion SA in wat1 mutant though overexpression of the bacterial SA hydroxylase gene NahG restored plant susceptibility to R. solanacearum (Denance et al., 2013). Mutation of EIN2, an important component in ET signal transduction, dramatically delayed bacterial wilt on Arabidopsis, which did not happen on etr1-3, ein4-1 and eni3-1, other ET insensitive mutants (Hirsch et al., 2002). In addition, while, the jar1-1 mutant -lacking the bioactive JA-IIe- shows the same sensitivity to R. solanacearum as wild type plants (Hirsch et al., 2002), loss of function of the JA receptor COI1-1 enhances plant defense against to R. solanacearum (Hernandez-Blanco et al., 2007).

WRKY transcription factors, critical players in modulating plant resistance to
 phytopathogens, were also reported to function in plant defense to *R*.
 solancearum. WRKY27 mutation delays disease symptom development by
 modulating signaling between the phloem and the xylem (Mukhtar et al., 2008).
 Inactivation of WRKY53 also reduces wilt symptom caused by *R*.
 solanacearum (Hu et al., 2008).

In roots, the cell wall is the first physical layer of plant defense against pathogens. It is demonstrated that alteration of cell wall affects Arabidopsis defense to *R. solanacearum*. Cellulose synthases are required for secondary cell wall formation. Mutations of cellulose synthase genes (CESA4, CESA7 and CESA8) confer enhanced resistance to R. solanacearum independently of SA, JA and ET but dependent on ABA (Hernandez-Blanco et al., 2007). Similarly, the WALLS ARE THIN 1 (WAT1) gene is essential for secondary cell wall deposition. A mutation in WAT1 leads to reduced cell elongation and

 secondary wall thickness, but it also increases SA content and plant defense to
vascular *R. solanacearum* (Denance et al., 2013). Furthermore, pectin
homogalacturonan in the root cell wall was reported to be degraded after *R. solanacearum* infection (Digonnet et al., 2012).

Transcriptional profiles by RNA-seq have been employed to look for important events in plant defense against R. solanacearum in Arabidopsis. The *R.* solanacearum \triangle hrpB mutant has a dysfunctional TTSS and loses the ability to invade host plants (Vasse et al., 2000). Plants infected with this mutant exhibit increased plant defense to subsequent virulent strain infection. Microarray analysis of transcriptional changes in aerial part of plants treated with GMI1000 \triangle hrpB indicated that 26% of up-regulated genes were involved in the metabolism and signaling of ABA (Feng et al., 2012). In addition, comparison of transcriptional profiles from the aerial part of Arabidopsis Col-0 inoculated with GMI1000 at several time points identified many differentially expressed genes associated with ABA signaling pathways (Hu et al., 2008).

However, previous microarray studies focused on transcriptional changes in the aerial part of root-inoculated Arabidopsis with GMI1000. Since R. solanacearum is soil-borne and infects plant roots, direct investigation of transcriptional changes in infected plant roots at a series of time points will help disclosing the molecular mechanism of *R. solanacearum* infection. In this study, by means of high-resolution temporal analysis of host global transcriptional changes following pathogen infection, we identified several important events as the activation of the biosynthesis and signaling of different hormones, and further connected root structure changes to the transcriptional reprogramming following R. solanacearum infection. Our data provides a cornerstone to understand complicated regulation networks during the infection process of *R. solanacearum* in the root.

RESULTS

152 Characterization of Root Morphology Changes Following GMI1000 153 Infection

As previously reported, Arabidopsis seedlings roots exhibited primary root growth retardation, de novo root hair formation and cell death appearance around the root tip at 9 days after GMI1000 treatment (Lu et al., 2018). To refine the appearance time of the three root phenotypes, we investigated the root elongation of Arabidopsis seedling after infection with GMI1000 over time. Primary roots kept growing the first 24 hours post-inoculation (hpi). At 48 hpi, primary root growth was found to be inhibited by GMI1000 (Fig. 1A). Root hairs covered root tips around 24 hpi, while they did not appear in water-treated seedlings (Fig. 1B). Roots were immersed in Propidium lodide (PI) -a DNA/RNA dye used to investigate cell integrity- and observed under confocal microscope. Cells in the root meristem area were alive at 24 hpi but already dead at 48 hpi (Fig. 1C). In addition, lateral roots emerged from primary roots treated with GMI1000 at 72 hpi and became apparent at 96 hpi. The number of these secondary roots on Arabidopsis root treated with GMI1000 was 4-5 fold higher than in water-treated plants (Fig. 1A and Fig. S1). According to these root structure changes over time, we divided the initial root infection by R. solanacearum into four phenotypic stages: No symptoms (NS) stage at 0-12 hpi, Root Hair (RH) emergence stage at 12-24 hpi, Primary root growth arrest and Cell death (PC) stage at 24-48 hpi and Lateral Root (LR) emergence stage at 48-72 hpi.

175 Time Series of Global Transcriptional Re-Programming in Roots 176 Challenged with GMI1000

To understand the events taking place at different infection stages of *R*. *solanacearum*, we infected 7-day-old seedling roots *in vitro*, and collected root samples at 0 hpi, 6 hpi, 12 hpi, 24 hpi, 48 hpi and 96 hpi, extracted total RNA and sequenced the global transcripts of GMI1000-infected roots. Around 600

seedling roots were pooled into one sample. Three biological replicates per time point were directly subjected to RNA-Illumina sequencing. An average of 33.9 million clean reads (range from 26.9-41.5 million) with Q30 > 90% were obtained per sample. More than 94 percent of clean reads were mapped to the Arabidopsis genome (Table S1). Aiming to disclose the molecular mechanism of early infection process of R. solancearum, we respectively compared R. solancearum-infected root transcriptomes at different infection time points with those obtained in water-treated roots after 96h and in GMI1000-treated roots at time 0h. The time series expression profiles identified a total of 2698 Arabidopsis genes as differentially expressed genes (DEGs) based on their significance in fold-change expression (padj<0.05) and at least a two-fold change in expression level (-1>log2>1) (Fig. 2 and Supplemental Data Set 1).

To analyze the overall patterns in gene expression during *R. solancearum* infection, the 2698 DEGs were clustered into 11 hierarchical clusters based on their expression patterns over time (Fig. 2). The list of genes in each cluster is presented in Supplemental data Set 2. These clusters group sets of genes that were sequentially induced upon pathogen challenge over time. The cluster VI genes started increasing at 12 hpi and peaked at HR stage (24 hpi), then slowly dropped back to basal level, which was the most quick response to R. solanacearum infection. The maximum level of cluster IV and V genes was at RH stage and PC stage (48 hpi), later 12h than that of cluster VI. Then cluster V guickly decreased. Comparing with relative long-lasting expression pattern of cluster V and IV, the highest expression level of cluster III genes was more concentrated in PC stage. The cluster I and II genes went up to maximum level at LR stage (72 hpi) and 96 hpi, which are the last induction clusters. The down-regulated genes also showed temporoally modulated expression pattern. The earliest repressed-gene clusters are cluster VIII and XI, which happened at RH stage. Interestingly, unlike cluster VIII maintaining lower expression, a few genes in cluster XI suffered a second induction at LR stage. The lowest expression level of cluster IX and cluster X occured at LR

stage.The expression of cluster VII were inhibited at LR stage and 96 hpi(Fig.2).

To check whether the co-expressing genes in the same cluster participated in similar biological processes, we investigated over-representation of Gene Ontology (GO) terms in these groups. The selected over-represented GO terms are shown at the right of each gene expression cluster in figure 2. Cell wall organization genes enriched in cluster VI unregulated before the appearance of root hairs (12 hpi) and reached their highest level at RH stage in response to the pathogen, reflecting cell wall remodeling has a specific role in the plant response to GMI1000 infection. A significant GO term in cluster V was lignin metabolic process. Cluster IV contained major GO terms: tryptophan metabolic process, auxin metabolic process and glucosinolate biosynthetic process, which share major components in their biosynthesis and peak at RH and PC stage. The GO term "response to auxin" was over-presented in cluster I, cluster II and cluster III and strongly induced during PC stage and LR stage (48-72 hpi), later 24 hour than GO term "axuin metabolic process" in cluster VI (Fig. 2). Additionally GO terms such as "response to JA", "response to abiotic stress", "response to heat" and "response to hydrogen peroxide" were also overrepresented in clusters I, II and III. GO terms related with plant defense such as "response to chitin", "response to bacterium", "response to SA" and "defense response" were enriched in cluster VII, which were significantly suppressed during LR stage (72 hpi) . Interestingly, GO terms "cell to cell junction" and "cell wall organization" also were enriched in Cluster VIII were significantly suppressed at PC and LR stage. Cluster X genes were significantly related with GO term "root hair cell differentiation", which suppressed when lateral root emerging.

Biological processes that take place during *R. solanacearum* infection are likely to affect the outcome of the plant-pathogen interaction. Therefore we further investigated enriched GO terms in the DEGs at single time points irrespective of the previous clustering (Fig. S2). This analysis revealed that cell

wall organization-associated genes were enriched at NS and RH stages, suggesting that these genes probably contribute to loosening the cell wall and cell-to-cell junctions, which may help R. solanacerum crossing the cortex and endodermis at early infection stages. The term "tryptophan metabolic process" was overrepresented in up-regulated DEGs at RH stage, which may point at tryptophan as a likely substrate for auxin biosynthesis. "Response to biotic stimulus" was a GO term overrepresented in up-regulated DEGs at RH stage and PC stage. "Response to hormones" was overrepresented in genes specifically upregulated at PC and LR stages (48 hpi and 72 hpi, respectively), which may reflect the root structure changes that take place at the LR stage. The GO term "response to abiotic stimulus" was also highlighted in the upregulated DEGs at PC stage. The upregulated "Glucosinolate biosynthetic process" term spanned from LR stage to 96 hpi. JA is involved in root development and regulation of plant defense. The DEGs related to "Response to JA" term remarkably increased at 96 hpi. In down-regulated DEGs, the terms "transport", "cell wall organization" and "root development terms" were over-represented at PC and LR stage. These sequentially overrepresented GO terms during early R. solanacearum infection indicate that infection is a programmed dynamic event from the very beginning of the plant-pathogen interaction.

262 Ethylene-, Jasmonate-, Auxin- and Abscissic acid-dependent signalling 263 are altered following *R. solanacearum* Infection

The first and rate-limiting step in ethylene (ET) biosynthesis is the conversion of S-adenosyl Methionine to 1-aminocyclopropane-1-carboxylic acid (ACC) by ACC synthase (ACS). Five out of the nine *ACS* genes in the Arabidopsis genome (*ACS2*, *ACS6*, *ACS7*, *ACS8* and *ACS9*) were induced at PC stage. Interestingly, the expression of *ACS5* was inhibited at the same time (Fig. S3A). No ACC oxidase gene was identified in our RNA-seq data. However, its regulatror SHYG was induced at RH and PC stage (Rauf et al.,

2013) (Fig. S3A). Moreover, *ERF* transcriptional factors including *ORA59* and *ERF71* in response to ET were up-regulated or down-regulated following
GMI1000 infection (Fig. S3A). These findings suggest that ET biosynthesis
and signaling are involved in *R. solanacearum* infection.

The expression of several genes involved in jasmonic acid (JA) biosynthesis and degradation was also altered in our RNA-seg data. For example, LOX1 and LOX2, encoding 13-lipoxygenase were induced at PC stage. LOXs are responsible for converting α -linolenic acid to 13-hydroxyperoxy-octadecatrienoic acid (13-HPOT) in plastids, which is the first step in the production of the JA precursor (Wasternack and Hause, 2013). However, Acyl-coenzyme A oxidase (ACX4) and 3-ketoacyl-CoA thiolase (KAT5), which catalize JA biosynthesis from this precursor (Li et al., 2005) were repressed after inoculation. Three of the four Arabidopsis jasmonate-induced oxygenases (JAOs), which inactivate JA through hydroxylation (Caarls et al., 2017), were highly expressed in our data. Similarly, a hydroxyjasmonate sulfotransferase (ST2A) that inactivates JA functions (Gidda et al., 2003) was highly induced at LR stage (Fig. S3B). Jasmonate ZIM domain Proteins (JAZ1, JAI3, JAI5 and JAZ10), key negative regulators of JA signaling pathway, were strongly activated at LR stage and 96 hpi (Fig. S3B). In summary, the decrease in JA biosynthesis and increase in JA degradation and negative regulators suggests an inhibition of this pathway by R. solanacearum at late infection stages.

The components in auxin metabolism, auxin signaling and auxin transport were up-regulated from NS stage to LR stage (Fig. 3). TRP4, TRP5, TRP1, TRP3 and TSB2 encode five key components in the transformation of chorismate to the auxin precursor tryptophan (Zhao, 2010). All of them were up-regulated at RH stage (Fig 3A and Fig 3B). Members of two of the four tryptophan-dependent auxin synthesis pathways described in Arabidopsis (Zhao, 2010; Rosquete et al., 2012) were up-regulated at RH stage (genes CYP79B2, CYP79B3, NIT1, NIT3 and YUC9) (Fig. 3A and Fig. 3B). In addition,

the expression of DAO1 and DAO2 -encoding genes that oxidate IAA to oxIAA and GH3 family genes, which conjugate amino acids to IAA (Rosquete et al., 2012) were all induced at RH and PC stages (Fig. 3A and Fig. 3B). Accumulation of auxin-responsive transcripts such SAURs and Aux/IAAs was observed at PC stage (Woodward and Bartel, 2005), which is 24 hours later than the peak auxin synthesis genes (Fig. 3B and Fig. S4). The expression of auxin response factors such as Auxin Response Factor 4 (ARF4) increased during infection (Fig. 3C), as well as the expression of auxin efflux transporters (PINs and ABCB4) (Rosquete et al., 2012), which increased at PC and LR stages (Fig. 3D). Moreover, a few regulators of stability of auxin transporters (PATL2, RAM2, PBP1, PILS7, SMXL8 and PID) were also differentially expressed in our data.

Our RNA-seq data also identified a group of genes that were associated with abscissic acid (ABA) metabolism and signaling (Fig. 4A). The expression of CYP707A, which oxidizes and inactivates ABA (Saito et al., 2004), was induced at all time points after 24 hpi. Expression of the ABA receptor PYL5 was inhibited after infection (Fig. 4A). And the ABI2, HAB1 and PP2C5 genes, encoding protein phosphatases that suppress ABA signalling through dephosphorylation of SNRK2 proteins (Umezawa et al., 2009), were both up-regulated at PC stage. Expression of OST1, essential for ABA signaling (Fujii et al., 2009), showed a peak at PC stage, and then guickly decreased at LR stage. On the contrary, other SNRK family genes were inhibited at PC and LR stage (Fig. 4A). Finally, expression of the ABA-dependent transcription factor ABF2 (Fujita et al., 2005) peaked at RH stage, 24 hours earlier than OST1 (Fig. 4A).

327 ABA signaling is involved in plant resistance to *R. solanacearum*

Next, we investigated whether the alteration of ABA biosynthesis and signalling caused upon *R. solanacearum* infection had an impact on plant responses to this pathogen. To this end, we took advantage of available

Arabidopsis mutants: the quintuple pyl1/pyl2/pyl4/pyl5/pyl8 (12458) and the sextuple pyr1/pyl2/pyl4/pyl5/pyl8 (112458) mutants, which are devoid of multiple ABA receptors and show reduced vegetative growth and seed production (Gonzalez-Guzman et al., 2012). We grew the Col-0 accession and ABA receptor mutants and tested their sensitivities to R. solanacearum infection. Both mutant lines showed increased wilting symptoms at 15 days post-inoculation compared with their wild type counterpart (Fig. 4B). This was translated into a significantly higher plant mortality rates in the mutants than in wild type plants (Fig. 4C). These results indicate a role of ABA signaling in plant resistance to R. solanacearum. We further tested if ABA signaling could affect the previously-described root morphology changes induced by the bacterium. The sextuple mutant exhibited root morphogenetic responses identical to wild type plants (Fig. 5), suggesting that ABA signaling is not required for *R. solanacearum*-induced root structural changes.

Regulation of Plant Defense Response genes in *R. solanacerum*-infected

Roots

Among the 2698 genes differentially-expressed after R. solanacearum infection 109 have been previously involved in plant defense (Fig. S5 and Supplemental Data Set 3). RLK3, RD19 and WRKY27 regulate plant defense to *R. solanacearum*. RLK3 encoding a cysteine-rich repeat receptor like kinase was induced in the Arabidopsis ecotype Niederzenz (Nd-1) infected with R. solanacearum GMI1000 (Czernic et al., 1999). RLK3 was strongly induced at 12 hpi and reached a peak at PC stage in infected plants (Fig. S5). Surprisingly, RD19, a cysteine protease required for RRS1-R-mediated resistance to R. solanacearum (Bernoux et al., 2008) was strongly inhibited upon infection (Fig. S5). Similarly, WRKY27, which was shown to promote disease symptom development (Bernoux et al., 2008), was repressed upon infection (Fig. S5). The negative regulators of pathogen-associated molecular patterns (PAMP)-triggered immunity (PTI) PUB22 and PUB23 (Trujillo et al., 2008),

were differentially expressed with PUB22 downgoing and a earlier induction peak on PUB23. *LYK4* participating in sensing chitin was induced at 12h after infection and the expression of other PTI regulators (*PEP1*, *PUB23* and *MPK11*) was strongly induced at RH and LR stages. Interestingly, these genes were inhibited at LR stage (Fig. S5). Finally, 6 WRKY, 2 ERF and 2 ANAC transcription factors, key modulators of plant immunity, were also identified as DEGs in our experiments (Supplemental Data Set 3).

Transcriptional regulation of Programmed Plant Cell Death genes in *R. solanacearum*-infected roots

Programmed cell death (PCD) in root tip cells was initiated around 24 hpi and completed around 48 hpi after infection (Fig. 1C). In line with cell death appearance in root meristem zone, many regulators of plant cell death (PCD) were differentially expressed (Fig. 6). For instance, expression of two negative regulators of cell death -MC2 and SYP122- (Zhang et al., 2008) (Coll et al., 2010) were strongly inhibited by *R. solanacearum* from RH stage on (Fig. 6A). of Consistent with down-regulation SYP122. the expression of mono-oxygenase1 (FMO1), required for SYP22-dependent lesion formation reached a peak at 24 hours after infection (Fig. 6A). Auto-inhibited Ca²⁺-ATPase 4 (ACA4), also involved in regulation of PCD (Boursiac et al., 2010) was repressed at 24-48 hpi (Fig. 6A). In addition, we also noticed plant senescence genes associated with PCD differentially expressed (Fig. 6B). Oresara1 (ORE1), a transcription factor regulating ET-mediated age-induced cell death, and WRKY57, a negative regulator of JA-induced leaf senescence (Jiang et al., 2014) were both strongly induced at RH stage, the latter starting induction at 12 hpi and decreasing at PC stage (Fig. 6B).

388 Root Architecture Responses to *R. solanacearum* Infection

Root hair formation was induced at RH stage at the root tip (Fig. 1B). We thus scrutinized our transcriptomes for differentially-expressed genes

described in the literature to play a role in this process. We found the root hair initiation zinc finger protein 5 (ZFP5) (An et al., 2012) and the Oxidative signal-inducible 1 (OXI1) kinase required for normal root hair development (Rentel et al., 2004) were induced at 6hpi, peaking at RH stage and returning to basal levels at LR stage (Fig. 7A). The ERU, EXP7 and LRX1 genes, involved in root hair elongation (Baumberger et al., 2001; Lin et al., 2011; Schoenaers et al., 2018), were also guickly turned on at NS stage (6-12 hpi) and inactivated at PC stage (Fig. 7A). According to these data, root hairs should appear on root tips just after 12 hpi. We thus analysed in further detail root hair appearance by observing infected root tips at 6, 12, 18 and 24 hpi. Appearance of root hairs around the root tip was observed at around 18 hours after infection (Fig. 7B), which correlates to the changes in root hair gene expression patterns.

Another dramatic response to R. solanacearum infection is root growth inhibition. In our transcriptome data, many regulators involved in primary root growth were identified (Fig. 8). The expression of several negative regulators of root growth increased after infection, reaching the highest levels at PC stage. These included the CLV3/ESR-related peptide 20 (CLE20) (Meng and Feldman, 2010), the methyltransferase PXMT1 (Chung et al., 2016), the triterpene synthesis genes THAH1, THAD1 and THAS1 (Field and Osbourn, 2008), the LRP1 gene -involved in root growth retardation induced by phosphate deficiency (Svistoonoff et al., 2007) and EFR, whose gain-of-function mutant showed shorter primary roots in rice (Xiao et al., 2016). On the contrary, positive root growth regulators were repressed at PC stage. Amongst them are GA3ox catalyzing the final step in gibberellic acid (GA) biosynthesis (Mitchum et al., 2006) and CLE6, whose overexpression in a ga3ox mutant partially restored primary root growth (Bidadi et al., 2014). Therefore, coordinated expression of positive and negative regulators may control root growth inhibition induced by *R. solanacearum*.

420 The last morphogenetic change observed in infected roots was enhanced

appearance of secondary roots at 72 hpi. The transcript levels of the lateral root formation repressors CLE1, CLE3 and GLIP2 (Lee et al., 2009; Araya et al., 2014) were significantly decreased during LR stage. In addition, the positive secondary root regulators GATA23, were induced at 24 hpi and repressed from 48 to 96 hpi (Fig. 8). Interestingly, the action of GATA23 is auxin-mediated (Xie et al., 2000; Lally et al., 2001; De Rybel et al., 2010; Lee and Kim, 2013), which suggests that auxin may be controlling this root response to *R. solanacearum*.

DISCUSSION

R. solanacearum causes genome-wide transcriptional reprogramming in 432 Arabidopsis

Transcriptional reprogramming in aboveground tissue following soil-drenched R. solanacearum has been previously reported in Arabidopsis (Hu et al., 2008; Feng et al., 2012). Leaf transcriptome analysis from susceptible plants showed that 40% of the up-regulated genes were involved in ABA biosynthesis and signaling (Hu et al., 2008), which is line with our root transcriptome results. Similarly Feng and colleagues found that 26% of the upregulated genes in the leaf transcriptome pretreated with nonpathogenic *Ralstonia* strain were also involved in ABA biosynthesis and signaling. These indicate ABA signaling is triggered by pathogenic and nonpathogenic invasion and may function in root defense agnist R. solanacearum. Very few SA-associated genes were found in our root transcriptome, which also happened in the leaf transcrptome (Hu et al., 2008). This corroborates the notion that SA does not have a key role in plant defense responses against many root pathogenic bacteria. Moreover, several genes involved in auxin signaling were down-regulated in the leaf transcriptome (Hu et al., 2008). In contrast, the auxin biosynthesis, signaling and transport pathways were significantly induced in the root transcriptome reported here. This discrepancy in the results could be partly caused by the different tissues used in the

experiment (leaf vs. root) and different inoculation methods employed (soil
drench vs. *in vitro* infection).

R. solanacearum manipulates different plant hormonal pathways

Plant hormones are well-known to synergistically or antagonistically affect each other's output, leading to plant resistance or susceptibility to various pathogens (Berens et al., 2017). Therefore, phytopathogens have acquired the abilities to hijack plant hormones to promote their proliferation in the host (Ma and Ma, 2016). Ethlyene and Jasmonic acid signals have been shown to be the main target of many virulence factors produced by biotrophic and hemibiotrophic phytopathogen, due to their negative role in plant immunity against biotrophic pathogens via SA antagonism (Kloek et al., 2001; Berrocal-Lobo et al., 2002). Ethlyene is produced by many plant pathogens including the bacterial pathogen Pseudomonas syringae and R. solanacearum (Weingart and Volksch, 1997; Valls et al., 2006). Disruption of ET production affects the virulence of *P. syringae* on soybean and bean (Weingart et al., 2001). In R. solanacearum, mutation of ethylene-forming enzyme (RsEFE) did not affect its proliferation on plant host (Valls et al., 2006). However plants defective in ethylene signaling (ein2 mutants), show delayed wilt symptom (Hirsch et al., 2002). Our transcriptome data shows that R. solanacearum highly induces expression of ACS genes in the roots, which could indicate that besides directly producing ET, R. solanacearum employ another unknown stratedy to activate endogenous ET.

The *P. syringae* virulence factors Coronatine, HopZ1a, HopX1 and AvrB, virulence factors, activate JA signaling by promoting degradation of JAZ proteins, key negative regulators in JA signaling (Melotto et al., 2006; Jiang et al., 2013; Gimenez-Ibanez et al., 2014; Zhou et al., 2015). Activation of JA signaling leads to entry of phytopathgoen into apoplast by reopening closed stomata and attenuate SA-dependent plant defense (Melotto et al., 2006; Zhou et al., 2015). Hernandez-Blanco reported mutation in JA-Ile receptor gene,

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Coronatine- insentistive 1 (COI1), conferred plant resistance to R. solanacearum (Hernandez-Blanco et al., 2007). Our data showed JA biosynthesis and degradation genes (LOX1, LOX4 and KAT5) were differentially-expressed at earlier RH stage and JAZs were mainly induced at LR stage and 96 hpi, suggesting that JA signaing pathway was activated and then qucikly inhibited during R. solanacearum infection. However, the jai3-1, *jar1-1*, and *dde2* mutants with disabled JA biosynthesis or signaling showed similar root architectures as wild type plants in response to this pathogen (Lu et al., 2018). This indicates that JA may be involved in plant defense, but not in the root morphogenesis changes caused by *R. solanacearum*.

Auxin signaling and transport has been reported to be manupilated by phytopathogens to suppress activation of SA-dependent defense. The P. syringae effector AvrRpt2 activates auxin biosynthesis and induces expression of auxin-response genes by promoting degradation of the key negative regulators of auxin signaling AUX/IAAs. The effector HopM1 also from P. syringae and PSE1 from *Phytophthora parasitica* disrupt auxin transport by affecting expression or localization of different PIN auxin transporters, which promotes pathogen infection by antagonizing SA signaling (Nomura et al., 2006; Chen et al., 2007; Cui et al., 2013; Evangelisti et al., 2013; Tanaka et al., 2013). Many plant pathogens, including R. solanacearum, produce auxin-like molecules, which may alter auxin homestasis and affect auxin signaling in the host plants (Manulis et al., 1994; Glickmann et al., 1998; Valls et al., 2006; Robert-Seilaniantz et al., 2007). Interestingly, we oberserved that auxin biosynthesis genes were activated at the RH stage by R. solanacearum . Auxin signaling and transport were also upregulated at PC and LR stage. In line with our data, The expression of *DR5*, a marker gene of auxin signaling pathway, was strongly induced in root vascular after R. solanacearum GMI1000 infection (Lu et al., 2018). Moreover, the *dgl1-1* tomato mutant with disordered auxin transport was found to be highly resistant to R. solanacearum (French et al., 2018). Together, these data strongly supports the notion that auxin

signaling plays a negative role in root defense against *R. solanacerum*. A
deeper understanding of the role of auxin signaling in plant susceptibility to *R. solanacearum* awaits further investigation.

ABA also plays an important role in attenuating plant defense, possibly by inhibiting SA signaling (Cao et al., 2011). Increase of ABA levels in infected plants will enhance plant susceptibility to the bacterial pathogen P. syringae, the fungus Magnaporthe grisea and the nematode Hirshcmaniella oryzae (de Torres-Zabala et al., 2007; Jiang et al., 2010; Nahar et al., 2012). In turn, various pathogenic fungi have been shown to produce ABA (Ma and Ma, 2016) and the effectors AvrPtoB and HopAM1 produced by P. syringae enhance plant susceptibility to the bacterial infection by promoting ABA biosynthesis or affecting ABA singaling (de Torres-Zabala et al., 2007; Goel et al., 2008). ABA also can positively regulate plant denfense to *P. syringae*. For example, ABA induces stomata closure and locks pathogen outside of host upon encountering pathogen, protecting plant from pathogen infection (Melotto et al., 2006). A large number of ABA-responsive genes were up-regulated in plants infected with the non-virulent R. solanacerum mutant $\Delta hrpB$ and in R. CESA4/CESA7/CESA8-mediated resistance to solanacearum (Hernandez-Blanco et al., 2007; Feng et al., 2012). abi1-1 and abi2-1, two ABA-insensitive mutants, exhibited more sensitivity to R. solanacearum and disabled *A* hrpB-triggered and CESA4/CESA7/CESA8- mediated plant resistance (Hu et al., 2008). Here we show that ABA signaling in root is turned on at PC stage, much earlier than activation of ABA signaling in leaf. Further genetic analysis demonstrated that simultaneous disruption of ABA receptors (12458 and 112458) dramatically enhanced susceptibility towards R. solanacerum. Consistent with this result, most components of the ABA receptor, PYR1, PYL1, PYL2, PYL4, and PYL8, express in the stele of root (Gonzalez-Guzman et al., 2012; Antoni et al., 2013). Interestingly, although both ABA receptor mutants are insensitive to ABA-mediated root growth inhibition (Antoni et al., 2013), they are still sensitive to root hair formation, root

541 growth inhibition, and lateral root formation caused by *R. solanacearum*. This 542 indicates that ABA signaling is not essential for *R. solanacearum*-triggered root 543 architechture changes. Together, our data suggests that ABA has a positive 544 effect on plant defense against *R. solanacerum*. However, the precise 545 mechanism by which ABA promotes defense to this bacteria still needs to be 546 further elucidated

Together, all these data indicates that the interplay between R. solanacearum and Arabidopsis is mediated by a complex interplay of hormones. In particular, aa synergistic effect among JA, ET, SA, ABA, and auxin seem to determine the level of defense to *R. solanacearum* in the plant in spatiotemporal way. Our data provides new insight into the signaling network that occurs in the root host in response to a root pathogen.

R. solanaceaum infection triggers specific defense responses in the root PTI and Effector-triggered immunity (ETI) are the two layers of defense that plant pose to phytopathogens (Jones and Dangl, 2006). In our RNA-seg data, we identified several components of both defense branches, which is consistent with the reports that PAMPs elicits transcriptional changes and callose deposition in Arabidopsis root and the effector RBP1 from root nematode Globodera pallida triggers Gpa2-dependent resistance and cell death (Sacco et al., 2009; Millet et al., 2010). LYK4, PUB22, PUB23 and, PEP1 and MPK11, components of PTI signaling are guickly induced upon infection. Interestingly, all of these PTI-related genes were inhibited at LR stage, suggesting that *R. solanacearum* infection represses PTI in the root. In addition, we also found around 19 NBS-LRR resistance genes in DEGs including ZAR1. ZAR1 detects the acetylated hopz-ETI-deficient 1 (ZED1) by the *P. syringae* effector HopZ1a and triggers ETI (Lewis et al., 2010; Lewis et al., 2013). This suggests that this NB-LRR might be involved in R. solanacearum effector recognition.

Accompanying with ETI, hypersensitive response (HR), a local cell death

at the attempted entry site of pathogens, often happens. Cell death was observed on root tips at PC stage after R. solanacearum infection. Interestingly, the occurrence of *R. solanacearum*-mediated cell death at the root tip is dependent on the presence of a functional type three secretion system (Lu et al., 2018). This could indicate that this cell death occurs via effector recognition and thus ETI would be occurring at R. solanacearum infecting roots. HR in leaf is thought to directly kill invaders and/or to interfere biotrophic pathogen with acquisition of nutrients (Heath, 2000). But we showed cell death in root seems not to affect the virulence of GMI1000 on Arabidopsis and we know GMI1000 is a compatible strain on Arabidopsis. Necrotrophic pathogen triggers cell death in order to obtain more nutrients that helps them accomplishing their life cycle (Glazebrook, 2005). Whether R. solanacearum would follow a similar strategy with the root tip or it is simply a consequence of infection needs to be answered.

Root morphgenesis changes triggered by *R. solanacearum* infection are accompanied by deep transcriptional reprogramming of genes involved in root architecture

The root is embedded in the soil and its architecture determines the efficiency for nutrient uptake and aboveground growth. Root architecture is often shaped by biotic stress and abiotic stress such as interaction with mutualist microbes and elements deficiency (Le Fevre et al., 2015). Several R. solanacearum strains cause root morphological changes (Lu et al., 2018), reminiscent of root morphological changes triggered by plant growth promoting bacteria/rizobacteria or fungi (PGPB/PGPR and PGPF) (Verbon and Liberman, 2016). These beneficial microbes affect cell division at the root meristem region and cell differentiation at sites of lateral root formation through manipulating endogenous hormone levels, hormone signaling such as auxin signaling and transports and metabolic processes, resulting in root structure changes (Verbon and Liberman, 2016). Our transcriptiomic analysis indicated

that auxin synthesis, signaling and transport in root are all activated by R. solanacearum colonization. The auxin insensitive single mutant tir1 and double mutant *tir1/afb2* were unable to form root hair in response to *R. solanacearum* infection (Lu et al., 2018). In consonance with this, IAA28 controlling the specification and identity of lateral root founder cells were upregualted in our data (De Rybel et al., 2010). This suggests that auxin signaling in relation to lateral formation might be activated in response to *R. solanacerum*. However, activation of auxin signaling enhances plant sensitivity to P. syringae, Xanthomonas oryzae, and Magnaporthe oryzae (Kazan and Lyons, 2014), whilst destruction of polar auxin transport in tomato tremendously elevated plant resistance towards R. solanacearum infection. This poses the question of whether the observed R. solanacearum-triggered architecture changes are side effects of elevated auxin levels caused by R. solanacearum to accomplish successful colonization or not. In addition, it is still not clear why R. solanacearum and PGPRs induce similar root architectures but exert two opposite influences on plant survival and what benefits does R. solanacearum obtain (if any) from altering root structure. Y.C.

MATERIALS AND METHODS

Plants Materials

In this study, Arabidopsis thaliana Col-0 and the ABA receptor mutants 12458 and 112458 were sown in soil and grown in the chamber at 23 °C, short day conditions(8h light, light intensity 12000 lux) and 70% humidity. For Arabidopsis seedling growth, Col-0 seeds were sterilized with 30% bleach and 0.02% TritonX-100, then sown on Murashige Skoog without sucrose (MS-) plate and grown with the plates set vertically at 25 °C and long day conditions (16h light, light intensity 9000 lux for 6-7 days).

R. solanacearum Infection.

The strain *R. solanacearum* GMI1000 was used to infection in this study. For soil drench infection assay, 5-week old plants were watered with a suspension

> of 1x10⁸ colony forming units (cfu). One hour later, roots of the infected plants were wounded three times with a blade, then grown into the chamber at 25 °C, 16h light. Leaf wilting symptoms and the number of dead plants were recorded over time. For *in vitro* infection we used the method previously described in Lu et al (Lu et al., 2018). Briefly, 6-7 day-old Arabidopsis seedlings grown on MS plates were inoculated 1cm away from root tip with a droplet of a solution containing 1x10⁷ cfu of *R. solanacearum* GMI1000, then kept into the growth chamber as detailed above. Root structures were photographed at indicated time points with an Olympus SZX16 microscopy and lateral roots were counted at the indicated times. For the cell death assay, seedlings were immersed into 0.1mg/ml propidium iodide solution and observed under an Olympus confocal microscope IX83-FV1200.

Sample Preparations for RNA-seq

The root samples were collected from around 600 infected seedlings at the indicated time point and frozen in liquid nitrogen, then directly sent to Novogene Company (Beijing, China) which performed RNA seg and data analysis (Supplemental method)

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10 9 11	931	syringae Effector Protein AvrB for Stomatal Invasion. The Plant cell 27:2032-2041.
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13 9	933	SUPPORTING INFORMATION LEGENDS
14 15 9	934	Figure S1. GMI1000 promotes lateral root formation. 7-day-old Arabidopsis
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17 9	935	seeding roots were inoculated with GMI1000 suspension or water. Lateral
18 19 9	936	roots were counted and recorded at 5 dpi.
20 21 9	937	Figure S2. Selected GO term overrepresented in differentially expressed
21 22 o	סכר	genes at different infection stages of CMI1000, CO terms in upper boxes
23	750	genes at unerent intection stages of OwnTood. OO terms in upper boxes
24 9	939	indicate up-regulated genes and GO terms in lower boxes indicate
25 26 9	940	down-regulated genes.
27		Figure 62 Discurthesis and simpling components of FT and 14 during the
28 9	941	Figure 53. Biosynthesis and signaling components of ET and JA during the
29 30 9	942	early stages of GMI1000 infection. (A) Heat map representation of differentially
31 9	943	expressed genes in ET biosynthesis and signaling pathways. (B) Heat map
32 33 g	944	representation of differentially expressed genes involved in JA biosynthesis
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35 9 36	945	and signaling pathways. The heat map depicts FPKM values after log10
37 9	946	transformation.
38	747	Figure S4 Activation of auxin nathway in response to GMI1000 infection. Heat
39 9 40	947	rigure 54. Activation of auxili pathway in response to Givin out infection. Heat
41 9	948	map values represent log ₁₀ -transformed FPKM values.
42 9	949	Figure S5. Transcriptional changes of part of differentially expressed genes
43 44 a	250	involved in plant immunity. Heat man values represent log ₄₀ -transformed
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46 9	951	FPKM values.
47 48 9	952	Table S1. Overview of quality of RNA-seq data.
49	152	Data S1 EDKM values of 2609 differentially expressed gappe in root at each
50 9	153	Data SI. FFRM values of 2090 differentially expressed genes in root at each
52 9	954	time points after GMI1000 treatment
53 9	955	Data S2. Membership of 11 gene clusters.
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957 in plant immunity.

959 FIGURE LEGENDS

Figure 1. Time series of root structure after GMI1000 infection. (A) Root growth was recorded and digital images were taken images at indicated time points. The arrow indicates lateral roots. Dashed line indicates root growth arrest. (B) Root hair images were taken with an OLYMPUS SZX16 microscope at the indicated time points. (C) Cell death on the root tip was stained with a PI solution and images were directly taken with an Olympus confocal microscope. **Figure 2.** Clustering analysis of RNAseg data. The heat map represents the expression patterns of 2698 DEGs identified in our RNA-seq data. The vertical axis organizes genes according to co-expression patterns. The horizontal axis displays time points. Red represents genes with high expression while blue represents genes with low expression. The selected overpresented GO terms in each cluster were shown on the left. The heat map depicts FPKM value after log₁₀ transformation.

Figure 3. Expression patterns of part of genes related with auxin biosynthesis, signaling and transport. (A) Auxin biosynthesis processes and metabolic processes. Differentially expressed enzymes in our RNA-seq data are shown in black bold, otherwise enzymes are shown in gray bold. (B) Expression patterns of differentially expressed auxin biosynthetic genes in response to GMI1000 infection. (C) Expression patterns of differentially expressed auxin signaling components in response to GMI1000. (D) Expression patterns of differentially expressed auxin transport in response to GMI1000. The heat map depicts FPKM values after log₁₀ transformation.

Figure 4. ABA receptor mutants *12458* and *112458* showed more sensitivity to
GMI1000. (A) Temporal dynamics of ABA signal components after GMI1000
treatment. Heat map depicts the expression patterns of differentially expressed
ABA-responsive genes. (B) Wilt symptoms weredigitally imaged at 15 dpi. (C)
Mortality rate of the infected plants was recorded at indicated times. **

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987 indicates P<0.001 (Student's test) with respect to Col-0.

Figure 5. Mutations in ABA receptors did not abolish root architecture changes caused by GMI1000. (A) Inhibition of *112458* root growth. Primary root elongation length after infection was measured at 4 dpi. (B) Root hair formation on *112458* root tips. The images were taken with Olympus microscope. (C) Lateral roots on *112458* root. Lateral roots per seedling were counted and recorded at 4 dpi

Figure 6. Expression dynamics of components of programmed cell death over the infection time. (A) Heat map depicting differentially expressed genes in effector –triggered hypersensitive responses. (B) Heat map representation of differentially expressed components of senescence. Heat map values represent log₁₀-transformed FPKM values.

Figure 7. Expression of genes regulating root hair formation correlated with root hair formation. (A) Heat map representation of differentially expressed genes in root hair formation after GMI1000 infection. Heat map values represent log₁₀-transformed FPKM values. (B) Root hair appeared at 18h after GMI1000 infection. The pictures were taken with an Olympus microscope at the indicated time after infection.

Figure 8. Transcriptional dynamic changes of differentially expressed genes in
 root architecture. Heat map values represent log₁₀-transformed FPKM values.



Time series of root structure after GMI1000 infection

148x128mm (600 x 600 DPI)



Clustering analysis of RNAseq data

174x179mm (600 x 600 DPI)





221x315mm (600 x 600 DPI)



ABA receptor mutants 12458 and 112458 showed more sensitivity to GMI1000

223x624mm (600 x 600 DPI)





168x377mm (600 x 600 DPI)



Expression dynamics of components of programmed cell death over the infection time

123x91mm (600 x 600 DPI)



Expression of genes regulating root hair formation correlated with root hair formation

130x100mm (600 x 600 DPI)

PC LR

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KCS2

LACS2 LPR1

AIR12

THAD1 THAH1

THAS1

SUR1 HOS3

PXMT1

CLE20 ROSY1

ARK2

MLO11 ANR1

HRS1

LAC2 SCM

ARSK1

NIA2 CLE44

WDL1

AIR3

CLE41 PERK4

XTH20

WOX11

BHLH129

CLE3 GA3OX1

RSA1

SLAH1

LRP1 NRT2. 1

ERF2 WVD2

SBT3. 5 DRO1

NS NS NS RH



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SUPPLEMENTAL DATA

Deep sequencing reveals that early reprogramming of Arabidopsis root transcriptomes upon *Ralstonia solanacearum* infection

Re Perez

Overview

Supplemental Methods

Supplemental References

Supplemental Tables: Table S1

Supplemental Methods

Sample Preparations for RNA-seq

The root samples were collected from around 600 infected seedlings at the indicated time point and fronzen in liquid nitrogen, then directly sent to Novogene Company (Beijing, China) and perform RNA seq and data analysis there.

RNA Extraction, Library Preparation and Sequencing (Novogene)

RNA were extracted using Trizol. After RNA extraction, RNA quality and quantitiy were assessed with following equipments: Nano Photometer spectrophotometer (IMPLEN,CA, USA) for RNA purity, Qubit®RNA Assay Kit in Qubit® 2.0 Flurometer (Life Technologies, CA, USA) for RNA concentration, RNA Nano 6000 Assay Kit of the Bioanalyzer 2100 system (Agilent Technologies, CA, USA) for RNA integrity.

A total amount of 3 µg RNA per sample was used as input material for the RNA sample preparations. Sequencing libraries were generated using NEBNext® Ultra™ RNA Library Prep Kit for Illumina® (NEB, USA) following manufacturer's recommendations and index codes were added to attribute sequences to each sample. Briefly, mRNA was purified from total RNA using poly-T oligo-attached magnetic beads. Fragmentation was carried out using divalent cations under elevated temperature in NEB Next First Strand Synthesis Reaction Buffer (5X). First strand cDNA was synthesized using random hexamer primer and M-MuLV Reverse Transcriptase (RNaseH). Second strand cDNA synthesis was subsequently performed using DNA Polymerase I and RNase H. Remaining overhangs were converted into blunt ends via exonuclease/polymerase activities. After adenylation of 3' ends of DNA fragments, NEB Next Adaptor with hairpin loop structure were ligated to prepare for hybridization. In order to select cDNA fragments of preferentially 150~200 bp in length, the library fragments were purified with AMPure XP system (Beckman Coulter, Beverly, USA). Then 3 µl USER Enzyme (NEB, USA) was used with size-selected, adaptor-ligated cDNA at 37°C for 15 min followed by 5 min at 95 °C before PCR. Then PCR was performed with Phusion High-Fidelity DNA polymerase, Universal PCR primers and Index (X) Primer. At last, PCR products were purified (AMPure XP system) and library quality was assessed on the Agilent Bioanalyzer 2100 system

The clustering of the index-coded samples was performed on a cBot Cluster Generation System using TruSeq PE Cluster Kit v3-cBot-HS (Illumia) according to the manufacturer's instructions. After cluster generation, the library preparations were sequenced on an Illumina Hiseq platform and 125bp/150 bp paired-end reads were generated.

Data Analysis: Quality Control, Reads Mapping to Genome, Quantification of Gene Expression level and Differential Expression Analysis (Novogene)

Raw data (raw reads) of fastq format were firstly processed through in-house perl scripts. In this step, clean data (clean reads) were obtained by removing reads containing adapter, reads containing ploy-N and low quality reads from raw data. At the same time, Q20, Q30 and GC content of the clean data were calculated. All the downstream analyses were based on the clean data with high quality.

Arabidopsis genome and gene model annotation files were downloaded from Ensemble database Version 34 directly. Index of the reference genome was built using Hisat2 v2.0.4 and paired-end clean reads were aligned to the reference genome using Hisat2 v2.0.4 (Kim et al., 2015). We selected Hisat2 as the mapping tool for that Hisat2 can generate a database of splice junctions based on the gene model annotation file and thus a better mapping result than other non-splice mapping tools.

HTSeq v0.6.1was used to count the reads numbers mapped to each gene (Anders et al., 2015). And then FPKM of each gene was calculated based on the length of the gene and reads count mapped to this gene. FPKM, expected number of Fragments Per Kilobase of transcript sequence per Millions base pairs sequenced, considers the effect of sequencing depth and gene length for

the reads count at the same time, and is currently the most commonly used method for estimating gene expression levels (Trapnell et al., 2010).

Differential expression analysis of two conditions/groups (two biological replicates per condition) was performed using the DESeq R package (1.18.0) (Anders and Huber 2012). DESeq provide statistical routines for determining differential expression in digital gene expression data using a model based on the negative binomial distribution (Anders and Huber, 2010). The resulting P-values were adjusted using the Benjamini and Hochberg's approach for controlling the false discovery rate. Genes with an adjusted P-value <0.05 found by DESeq and fold change (-1>log2>1) were assigned as differentially expressed.

Clustering of Gene Expression Profiles (Novogene)

Cluster analysis is used to determine the expression patterns of differential genes under different experimental conditions. Heatmap represents the expressions of all differentially expressed genes identified in the RNA-seq experiment. The FPKM value of differential genes under different experimental conditions was used as the expression level, and hierarchical clustering analysis was performed. Different colored regions represent different clustering grouping information. The X-axis represents sample name and the Y-axis represents the differentially expressed genes. Expression data was normalized in log10 (FPKM+1) manner, heatmaps were drawn by R pheatmap package (Kolde, 2018).

GO Analysis

GO annotation analysis was performed using the Agrigo v2.0 (Tian *et al.*, 2017). Overrepresented GO_Biological_Process categories were identified using a hypergeometric test with FDR<0.05 with the whole annotated genome as the reference test.

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Supplemental Tables

Supplemental Table S1. Overview of quality of RNA-seq data.

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Sample name	Raw reads	Clean reads	Clean bases (Gb)	Mapped reads (% of total)	Q30(%)		
G0	34305303	33278781	4.99	31560849 (94.86%)	91.37		
G6	36097955	34998502	5.25	33069200 (94.50%)	92.30		
G12	39001013	37994586	5.70	36226397 (95.38%)	91.40		
G24	34175878	33150365	4.97	31537042 (95.16%)	91.74		
G48	33370411	32175593	4.82	30628497 (95.19%)	91.78		
G72	32732665	31887137	4.78	30067895 (94.51%)	91.44		
G96	35598467	34544686	5.18	32460784 (94.07%)	92.06		
H96	35542801	34630479	5.19	32848338 (94.89%)	91.74		







158x147mm (600 x 600 DPI)



PIN5 SAUR49 CYP83A1

0BP2 SAUR41 PIN6 ATSG13350 ARF5/IAA24 SAUR71 LRP1 PIN4 FRY1 SUR1 MES16 DA02 TRP5 TRP3 TRP3 TRP3 TRP1 TRP3 CVP79B2 TRP4 SAUR30 SAUR76 GH3.2 UGT74E2 UGT74E2 UGT74E2 UGT74E2 UGT74E2 IAA2 AUF1 SMXL8 DA01 AAGOS ZFP1 PID A8C64 GH3.3 IAA30

IAA9 At5g13370 PBP1 SAUR34 IAGLU IAA31 NIT3 IAA29 PILS7 AIR3 SAUR67 AR2 HCT GH3.6 SAUR66 SAUR66 AIR12 CYP59B3 CYP59B1

GSTUS AT4G18270 IAA28 UGT78D2 AIR1 GATA23 SAUR5 COBL9 SAUR72 SAUR75 SAUR75 SAUR75 SAUR75 SAUR76 SAUR76 SAUR78 SAUR76 SAUR32 GLIP2 SAUR40 RMA2

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72h

96h

24h

12h

23x40mm (600 x 600 DPI)



117x177mm (600 x 600 DPI)