

Salicylic Acid and Jasmonic Acid Pathways are Activated in Spatially Different Domains Around the Infection Site During Effector-Triggered Immunity in *Arabidopsis thaliana*

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The innate immune response is, in the first place, elicited at the site of infection. Thus, the host response can be different among the infected cells and the cells surrounding them. Effector-triggered immunity (ETI), a form of innate immunity in plants, is triggered by specific recognition between pathogen effectors and their corresponding plant cytosolic immune receptors, resulting in rapid localized cell death known as hypersensitive response (HR). HR cell death is usually limited to a few cells at the infection site, and is surrounded by a few layers of cells massively expressing defense genes such as *Pathogenesis-Related Gene 1* (*PR1*). This virtually concentric pattern of the cellular responses in ETI is proposed to be regulated by a concentration gradient of salicylic acid (SA), a phytohormone accumulated around the infection site. Recent studies demonstrated that jasmonic acid (JA), another phytohormone known to be mutually antagonistic to SA in many cases, is also accumulated in and required for ETI, suggesting that ETI is a unique case. However, the molecular basis for this uniqueness remained largely to be solved. Here, we found that, using intravital time-lapse imaging, the JA signaling pathway is activated in the cells surrounding the central SA-active cells around the infection sites in *Arabidopsis thaliana*. This distinct spatial organization explains how these two phytohormone pathways in a mutually antagonistic relationship can be activated simultaneously during ETI. Our results re-emphasize that the spatial consideration is a key strategy to gain mechanistic insights into the apparently complex signaling cross-talk in immunity.

Keywords: *Arabidopsis thaliana* • Effector-triggered immunity • Jasmonic acid • Salicylic acid • *Pseudomonas syringae* pv. *tomato* DC3000 carrying *AvrRpt2* • Time-lapse imaging.

Abbreviations: Act2, Actin 2; Chl, chlorophyll; ETI, effector-triggered immunity; h.p.i., hours post-inoculation; HR, hypersensitive response; JA, jasmonic acid; MPK, mitogen-activated protein kinase; *pPR1*, *PR1* promoter; *PR1*, *Pathogenesis-Related Gene 1*; *Pst_a2*, *Pseudomonas syringae* pv. *tomato* DC3000 carrying the *AvrRpt2* effector; *pVSP1*, *VSP1* promoter; qRT-PCR, quantitative reverse transcription-PCR; ROI, region of

interest; *RPS2*, *Resistance to Pseudomonas syringae* 2; SA, salicylic acid; *SID2*, *Salicylic acid Induction-Deficient 2*; *VSP1*, *Vegetative Storage Protein 1*; YFP, yellow fluorescent protein; YFP-NLS, YFP fused to the nuclear localization signal.

Introduction

Innate immunity is activated upon perception of pathogen-derived molecules by the host cells. Effector-triggered immunity (ETI), a form of innate immunity in plants, is activated by specific recognition of pathogen effector activities by their corresponding plant cytosolic immune receptors (Dodds and Rathjen 2010). Activation of ETI leads to transcriptional up-regulation of defense-related genes and is often associated with rapid localized cell death at the site of infection, known as hypersensitive response (HR) (Dodds and Rathjen 2010). Thus, HR cell death lesion limited to a few cells in the vicinity of the infection site appears to be surrounded by a few layers of cells massively expressing a number of defense genes such as *Pathogenesis-Related Gene 1* (*PR1*) (Schmelzer et al. 1989, Ohshima et al. 1990, Enyedi et al. 1992, Murray et al. 2002).

The plant hormone salicylic acid (SA), which plays a key role in plant immunity against biotrophic pathogens, has been implicated in the formation of such a virtually concentric pattern of the cellular responses observed in HR (Enyedi et al. 1992, Dorey et al. 1997, Fu et al. 2012, Yan and Dong 2014). A recent study using *Arabidopsis thaliana* also proposed that a concentration gradient of SA formed around the pathogen infection site elicits different defense responses, such as cell death and *PR1* activation, in a dose-dependent manner (Fu et al. 2012). Thus, the French flag model, which explains various processes of development, might also be applicable to explain how plant cells acquire positional information upon a pathogen attack to produce the organized local immune responses around the infection site (Wolpert 1969, Enyedi et al. 1992, Dorey et al. 1997, Fu et al. 2012).

A number of studies have demonstrated that the SA pathway functions antagonistically against a signaling pathway controlled by jasmonic acid (JA), another plant hormone required for

immunity against necrotrophic pathogens as well as defense responses upon physical wounding (Glazebrook 2005, Vlot et al. 2009). Due to the mutually inhibitory relationship between the SA and JA pathways, plants are considered to select either pathway depending on lifestyles of invading pathogens (Glazebrook 2005). Several studies, however, reported that JA is accumulated massively when ETI is evoked (Kenton et al. 1999, Spoel et al. 2003). Moreover, the JA pathway was shown to contribute positively to ETI (Tsuda et al. 2009, Liu et al. 2016). These findings suggest that the SA and JA pathways are activated simultaneously in the same plants only in the case of ETI. Accordingly, the spatially controlled trade-off between the SA and JA pathways within a single leaf was not observed exceptionally in the case of ETI (Spoel et al. 2007). These findings that are apparently contradictory to the well-established antagonistic relationship between the SA and JA pathways might be explained by the characteristic compensatory signaling network structure governing ETI and/or the proposed unique interplay between SA and JA only found during ETI (Tsuda et al. 2009, Liu et al. 2016). Alternatively, it could be explained by simply introducing another layer of complexity, e.g. a spatiotemporal aspect, into the study of plant immunity. Indeed, our knowledge about the spatiotemporal dynamics of the plant immune response remains largely limited (Murray et al. 2002).

Here, we report the spatiotemporal dynamics of defense-related promoter activities during ETI in *A. thaliana*. We established a time-lapse imaging assay of defense-related promoter activities using a fluorescent reporter protein in non-detached plant leaves, which enabled us to capture spatiotemporal development of plant immune responses. Using the system, we detected the spatiotemporal dynamics of the promoter activities of *PR1*, a conventional marker of SA activity, as well as of *Vegetative Storage Protein 1* (*VSP1*), a JA marker, for 40 h after hand-infiltration of *Pseudomonas syringae* pv. *tomato* DC3000 carrying *AvrRpt2* (*Pst_a2*). Our imaging data indicate that the SA and JA pathways are activated in distinct concentric domains; inner SA and outer JA domains around the HR cell death area, which explains the previous observations on the SA–JA relationship without contradiction. Our results shed light on the importance of spatial consideration as a key strategy to gain mechanistic insights into the apparently complex signaling cross-talk in plant immunity.

Results

The use of a nuclear-localized fluorescent protein discriminates fluorescent-based promoter activity from autofluorescence in planta

Plant immune response involves a dynamic transcriptional reprogramming regulated by multiple transcription factors as well as phytohormones (Tsuda and Somssich 2015). Accumulating data sets of a number of transcriptome profiling studies provide us with the opportunities to capture information on the plant immune system, e.g. the network structures of the signaling cascades as well as the temporal dynamics (Mine et al. 2014). However, our knowledge about the spatial

dynamism of ETI still remains limited (Murray et al. 2002). In order to extend our understanding of the spatial aspect of the plant immune system, we generated transgenic Arabidopsis plants expressing yellow fluorescent protein (YFP) fused to the nuclear localization signal (YFP–NLS) under the control of the promoters of the defense marker genes (Kubo et al. 2005). First, we focused on the promoter activity of the *PR1* gene, a well-established marker gene for immunity controlled by SA (Vlot et al. 2009). Hand-infiltration of *A. thaliana* (ecotype Col-0) leaves with a dense suspension of *Pst_a2* results in *Resistance to Pseudomonas syringae* 2 (*RPS2*)-2- (*RPS2*) mediated ETI associated with HR cell death (Yu et al. 1993). Autofluorescence derived from plant chlorophylls (Chls) was utilized as a marker to detect living plant cells (Guadagno et al. 2017). Using a wide-field fluorescent stereomicroscope with our setting described in the Materials and Methods, we succeeded in visualizing the HR lesion by loss of Chl autofluorescence (Fig. 1A). The use of the characterized 4.5 kb upstream sequence of *PR1* reproduces the characteristic pattern of *PR1* promoter (*pPR1*) active cells surrounding the HR lesion 22 hours post-inoculation (h.p.i.) (Fig. 1A) (Lebel et al. 1998, Murray et al. 2002). Infiltration of the mock solution did not activate *pPR1* (Supplementary Fig. S1). The *pPR1* activity is relatively high around the mid-rib, as observed for ETI triggered by another *Pseudomonas* strain (Murray et al. 2002). Although *pPR1* activity has been extensively studied and characterized by various means such as the luciferase reporter, the use of YFP–NLS enabled us to detect every single cell, in which *pPR1* is activated, in the infiltrated leaves of the *pPR1* reporter (hereafter, *pPR1*-YFP-NLS) plants, providing information on the promoter activation pattern with cellular resolution when combined with appropriate microscopes (Fig. 1B) (Murray et al. 2002). Furthermore, NLS-mediated accumulation of YFP in the nuclei allows us to distinguish the *pPR1*-driven YFP signal from plant autofluorescence accumulated during immune responses (Fig. 1B) (Bennett et al. 1996). In a magnified confocal image of the HR cell death border, only several layers of cells from the border exhibit massive YFP accumulation in the nuclei, indicating high *pPR1* activity (Fig. 1B). The nuclear YFP signals are suddenly decreased outside of the cell layers showing the *pPR1* maxima surrounding the HR lesion (Fig. 1B). This observation indicates that *pPR1* activity is spatially regulated and strictly confined to the vicinity of the infection sites, which is in accordance with the published data of other defense-related genes activated in HR (Schmelzer et al. 1989, Ohshima et al. 1990). Thus, the promoter–YFP–NLS reporter transgenic plants are useful tools to give us a good spatial resolution in our understanding of ETI without any fixation or enzymatic reaction.

In toto live-imaging of a whole intact leaf undergoing ETI triggered by *RPS2* revealed a dynamic spatiotemporal pattern of *pPR1* activation

Biological events such as immunity and development progress over time. Thus, we combined the reporter plants with an automated fluorescent stereomicroscope, allowing us to

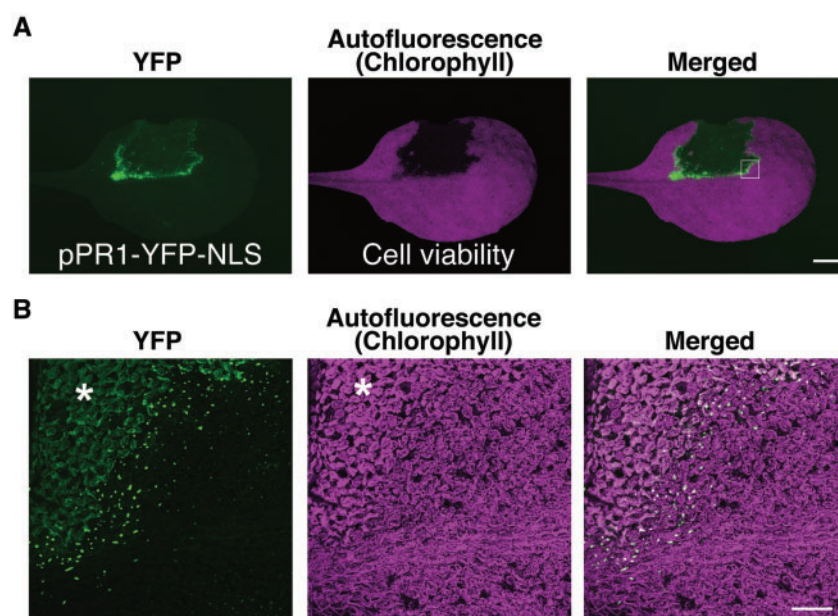


Fig. 1 The use of YFP–NLS to visualize in planta promoter activity. (A) Fluorescent stereomicroscopic images of a pPR1-YFP-NLS leaf partially infiltrated with *Pst_a2* ($OD_{600} = 0.2$) at 22 h.p.i. HR cell death was detected by the loss of Chl autofluorescence. Scale bar = 2.5 mm. (B) Confocal images of the region corresponding to the white square in (A). In our confocal setting described in the Materials and Methods, dead cells were visualized by whole-cell autofluorescence detected in both YFP and Chl autofluorescence images (in the area shown with an asterisk). They are distinguished from the nuclear- and chloroplast-localized signals. Scale bar = 200 μ m. (See also [Supplementary Fig. S1](#)).

perform intravital time-lapse imaging to capture the promoter dynamics in a soil-grown intact plant leaf over a few days. Since the light condition strikingly affected *PR1* accumulation during ETI, we designed the time-lapse program in which YFP and autofluorescence images were taken at 3 min intervals programmed to expose the plant specimen to the light (Zeier et al. 2004). Time-lapse imaging of a pPR1-YFP-NLS leaf infiltrated with a dense suspension of *Pst_a2* revealed that *pPR1* is transiently activated around the HR cell death lesion (**Fig. 2A**; [Supplementary Fig. S2A](#); [Supplementary Movie S1](#)). To analyze the temporal *pPR1* activation profile, we set multiple regions of interest (ROIs) containing a few cells to cover almost all of the *pPR1* active area at the margin of the HR lesion (white circles in **Fig. 2B**). The averaged data show an exponential increase of the YFP intensity, with a single peak around 12 h.p.i. followed by a gentle decrease by 40 h.p.i. (the white line in **Fig. 2C**). The YFP intensity profile after 24 h.p.i. could contain values derived from the background autofluorescence accumulated in the HR cell death area that cannot be distinguished from the YFP signal at this resolution (as in **Fig. 1**). Nevertheless, the averaged YFP intensity profile exhibits a transient activation of *pPR1* during ETI. Focusing on the YFP profiles of the individual single ROIs, not the averaged data, however, showed notable variations in the temporal intensity profiles, even in comparisons of two ROIs located close to each other (i.e. ROI 1 vs. ROI 2 and ROI 3 vs. ROI 4 in **Fig. 2B, C** and [Supplementary Movie S2](#)). This is also recognized as large SDs of the YFP intensity profiles during the *pPR1* active period (**Fig. 2C**). These tendencies of *pPR1* activation around the HR lesion described above are also observed reproducibly ([Supplementary Fig. S2B, C](#)). These observations imply that the kinetics of *pPR1*

activation at the cellular level are not as simple as one might expect from the analyses of bulked tissues.

As previously reported, we also observed sharp borders between the HR cell death area and the surrounding *pPR1* active area, with the remainder of the tissue showing no apparent response (**Fig. 2D**) (Schmelzer et al. 1989). Using 800 time-lapse images for 40 h after *Pst_a2* inoculation, we generated a Temporal-Color-Coded image of the YFP signals as well as a kymograph, both enabling us to capture the spatiotemporal dynamics of *pPR1* activation in *RPS2*-mediated ETI (**Fig. 2E, F**). The images reveal that the clear demarcation of the *pPR1* active zone was strictly maintained over 40 h around the lesion, indicating the presence of an active regulatory mechanisms to limit the propagation of the *pPR1* active area (**Fig. 2E, F**). In addition, we detected weak and transient *pPR1* activation in the uninfiltrated side of the leaf after the local *pPR1* activity declines (**Fig. 2E**; [Supplementary Fig. S2D, E](#)). Although the spatial pattern of this *pPR1* active area differs in every experiment, the weak *pPR1* activation in the uninfiltrated side of the inoculated leaves is reproducible ([Supplementary Fig. S2F](#)). Plants possess an ability to activate immunity systemically upon local pathogen challenges (Vlot et al. 2009). Our observation might indicate that a similar phenomenon happens even at the uninfected side of the inoculated leaf. Thus, our data obtained using intravital time-lapse imaging demonstrate that *pPR1* activation in *RPS2*-conditioned ETI appears to possess two modes of action; the primary strong activation around the infection sites within the first 24 h followed by the secondary weak activation in the uninfiltrated area. Spatial resolution achieved by imaging techniques enables us to detect, for the first time, these two phases of *pPR1* activation in spatially distinct area of leaves. Furthermore, it also reveals

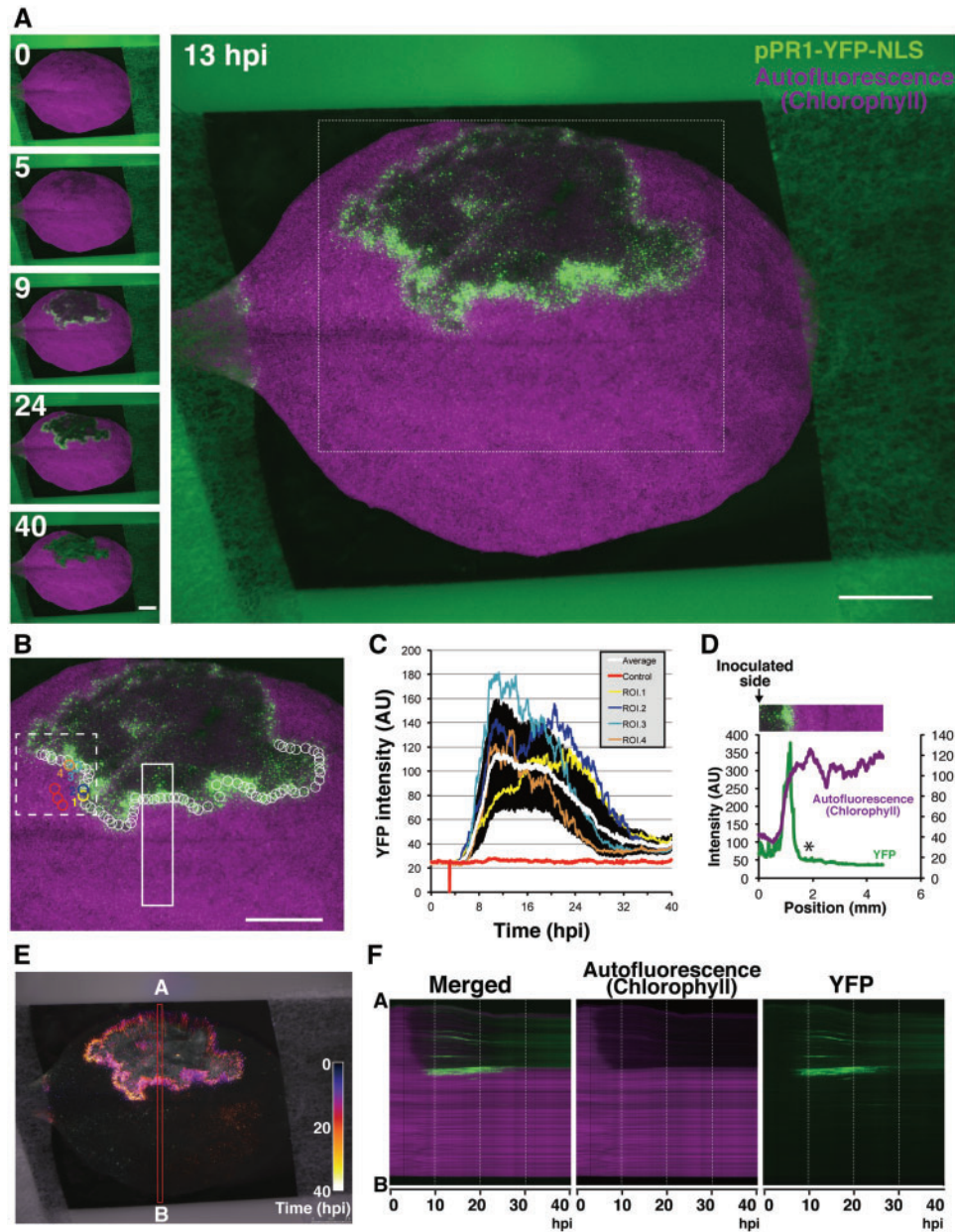


Fig. 2 Visualization of *pPR1* dynamics during *RPS2*-conditioned ETI. (A) Selected time-lapse images out of 800 images of the in vivo spatiotemporal dynamics of *pPR1* activity for 40 h after *Pst*_{a2} infiltration ($OD_{600} = 0.2$). Scale bars = 2.5 mm. (B) The region corresponding to the white box in (A) showing the positions of ROIs analyzed here. The red circles are used for background controls and the other circles are for calculating average intensity. The YFP intensity profiles in the numbered ROIs are individually shown in (C). The circles are all the same size. Scale bar = 2.5 mm. (C) YFP intensity plots in the ROIs shown in (B). Means \pm SD are plotted for the control ($n = 3$) and the others ($n = 70$). YFP profiles in four selected ROIs out of 70 ROIs are independently shown. The value 0 is due to the accidental loss of an image in the time-lapse system. (D) The intensity profiles of YFP and Chl autofluorescence in the white closed box shown in (B). A white asterisk indicates a sudden fall of YFP intensity outside the infection site. (E) Spatiotemporal dynamics of *pPR1* activity are shown in Temporal-Color Code. All the YFP images corresponding to (A) are re-colored by a specific Temporal-Color Code shown in the picture. (F) Kymographs corresponding to the red box along the A–B axis (E) are generated. (See also [Supplementary Fig. S2](#); [Supplementary Movies S1, S2](#)).

that individual cells exhibit temporal variation in *pPR1* activation, although the leaf taken as a whole appears to activate *pPR1* transiently with a single peak. Conventional molecular and biochemical techniques analyzing whole-leaf extracts might have overlooked such fluctuations in *pPR1* activation at the single-cell level.

The promoter of *VSP1*, a JA marker gene, is activated in the periphery of the *pPR1* active domain around the HR lesion

Recent studies showed a positive contribution of JA, in addition to SA, in invoking ETI conditioned by *RPS2* (Tsuda et al. 2009, Liu et al. 2016). JA and SA are known to function usually in an

antagonistic relationship, and thus simultaneous activation of SA and JA in ETI is considered to be a unique case (Vlot et al. 2009, Liu et al. 2016). However, those studies might have overlooked the spatial organization of two phytohormone signaling pathways. Therefore, we analyzed the activity of the promoter of *VSP1* (*pVSP1*), a JA marker gene, in response to infiltration of *Pst_a2* (Berger et al. 1995, Utsugi et al. 1998, Guerineau et al. 2003). Time-lapse imaging of a leaf of the transgenic plant carrying YFP–NLS under the control of *pVSP1* (*pVSP1*–YFP–NLS) detected transient and strong activation of *pVSP1* around the infection site (Fig. 3A; Supplementary Fig. S3A; Supplementary Movie S3). At the periphery of the infection site, *pVSP1* activity is observed earlier than that of *pPR1*, but activation kinetics of *pVSP1* show a variation to some extent (Supplementary Fig. S3B–F). Since a weak *pVSP1* activity is often observed along the vasculature, the position of the infection site in relation to the vascular system might affect the *pVSP1* activation kinetics during ETI (Supplementary Fig. S3A). After approximately 15 h.p.i., the *pVSP1* active area is gradually propagated from the infection site to the uninfected area, especially at the side of the leaf tip (Fig. 3A, B; Supplementary Fig. S3, AG; Supplementary Movie S3).

In contrast to the *pPR1* active domain observed just around the HR cell death border, a spatial gap was detected reproducibly between the *pVSP1* active domain and the HR cell death area despite a variation of *pVSP1* activation kinetics in ETI (Fig. 3C–E; Supplementary Fig. S3H). Mock treatment does not trigger such a pattern of *pVSP1* activation (Supplementary Fig. S3I). In addition, *Pst_a2*-infiltrated leaves without time-lapse imaging also produced a similar *pVSP1* activation pattern with a *pVSP1* inactive gap around the HR lesion (Supplementary Fig. S3J). These data suggest that this characteristic *pVSP1* activation pattern is not due to any stress from infiltration or continuous imaging. Since *pPR1* is activated just at the margin of the HR cell death domain, *pVSP1* activation is considered to occur mainly outside the *pPR1* active domain around the infection sites (Figs. 2D, F, 3C–E). These data strongly indicate that this virtually concentric pattern of the inner *pPR1* active and the outer *pVSP1* active domains around the infection foci is prominent in *RPS2*-triggered immunity.

SA accumulates in the *pPR1* active cells in early ETI

A recent study showed that, in ETI triggered by *RPS2*, *PR1* accumulation was maintained through MPK3/6 activity in the *salicylic acid induction-deficient 2* (*sid2*) mutants that fail to accumulate SA fully upon pathogen challenge, arguing that *PR1* might no longer be an SA marker in ETI (Tsuda et al. 2013). Therefore, we analyzed the spatial distribution of SA in relation to *pPR1* activity in leaves elicited by *RPS2*. Infiltration of one half of *pPR1*–YFP–NLS leaves with *Pst_a2*, not with the mock solution, resulted in YFP accumulation detectable only in the infiltrated side at 7 h.p.i. (Fig. 4A–C). Using these leaves, four different zones along the border between the infiltrated (zones 1 and 2) and uninfiltrated (zones 3 and 4) halves were analyzed for SA levels as well as endogenous *PR1* transcript

levels (Fig. 4A, D, E). As visualized by YFP, *Pst_a2*-infiltrated zones highly accumulate *PR1* transcripts (zones 1 and 2). The *PR1* level in zone 3 is only 5% of that in zone 2 in *Pst_a2*-treated leaves (Fig. 4D). A similar trend is observed for SA distribution. Zone 3 accumulates only 13% of the SA level in zone 2 in those leaves (Fig. 4D). Thus, in *RPS2*-conditioned ETI, massive SA accumulation largely coincides with high *PR1* transcript levels. An earlier study has shown that the transgenic plants constitutively expressing *NahG*, a bacterial SA-catabolizing enzyme, were unable to accumulate *PR1* transcripts in response to an incompatible pathogen (Delaney et al. 1994). Accordingly, the strong correlation between SA accumulation and the *PR1* transcript levels cannot be rejected in ETI in the wild-type background, although MPK3/6-mediated compensation of *PR1* activation might occur in the absence of *SID2* activity in ETI. Thus, the spatiotemporal pattern of *pPR1* activation observed in this study possesses a strong correlation with SA accumulation during ETI triggered by *RPS2*. Our data suggest formation of a steep SA concentration gradient around the infection site in early ETI (Fig. 4E), which is reflected as a sharp decrease of YFP intensity over the infiltrated/non-infiltrated border. Our data obtained in Arabidopsis agree with the previous findings in tobacco, suggesting conservation of a sharp SA gradient in ETI among plants (Enyedi et al. 1992).

As for JA, we detected higher JA accumulation from extracts of whole leaves challenged by *Pst_a2* at 24 h.p.i., compared with mock treatment (Supplementary Fig. S4A). Accordingly, the *VSP1* transcript levels are also found to be accumulated in the pathogen-challenged whole-leaf samples at 24 h.p.i. (Supplementary Fig. S4B). Using the same site-specific samples as used for *PR1* analysis, we examined the spatial pattern of *VSP1* accumulation (Supplementary Fig. S4C). Although the variance in the *VSP1* transcript levels in the respective samples is very high, as expected from live-imaging data, there appears to be a trend that the uninfiltrated area (zones 3 and 4) showed higher *VSP1* accumulation than the infiltrated area (zones 1 and 2) in the pathogen-treated leaves, supporting our finding that *VSP1* is activated outside the infected area where *PR1* is activated (Supplementary Fig. S4C). Thus, spatial separation of the *pPR1* and *pVSP1* active domains are confirmed, at least at the level of mRNA accumulation in the early *RPS2*-mediated immunity.

Discussion

Our data revealed that *PR1* and *VSP1* genes are activated in different domains, namely the local *PR1* and the peripheral *VSP1* domains, around the infection site where *RPS2*-conditioned immunity is triggered. *PR1* has been considered to be a conventional marker gene for SA in the case of the wild-type background (Vlot et al. 2009) and *PR1* transcript accumulation spatially correlates with SA accumulation in early *RPS2*-mediated immunity (Fig. 4A, D, E). *VSP1* is a well-established marker for a branch of the JA signaling pathway (Kazan and Manners 2013). Collectively, our finding indicates that the SA and JA pathways are spatially separated domains as exemplified

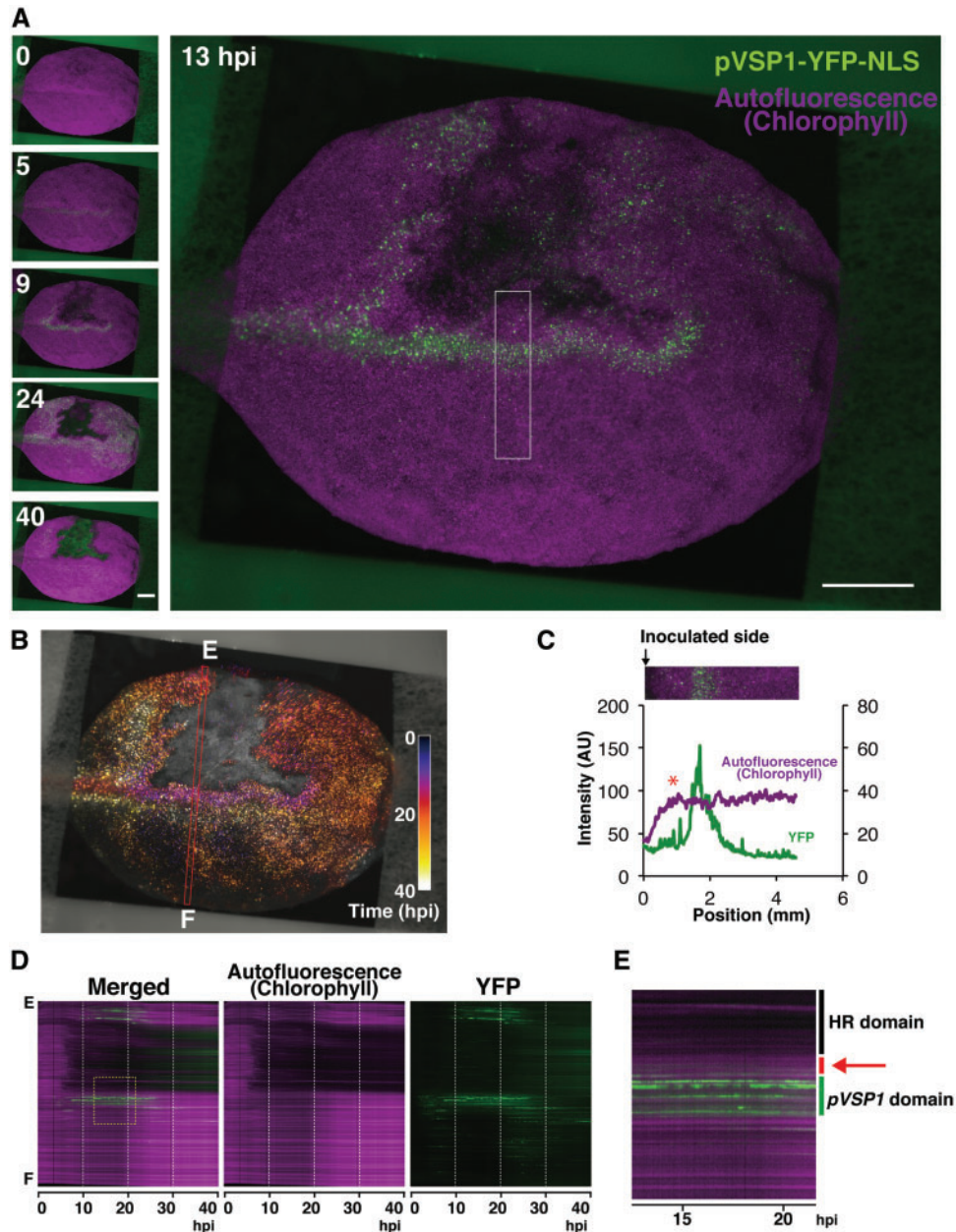


Fig. 3 Visualization of *pVSP1* dynamics during *RPS2*-conditioned ETI. (A) Selected time-lapse images out of 800 images of in vivo spatiotemporal dynamics of *pVSP1* activity for 40 h after *Pst*_{α2} infiltration (OD₆₀₀ = 0.2). Scale bars = 2.5 mm. (B) Spatiotemporal dynamics of *pVSP1* activity are shown in Temporal-Color Code. All the YFP images corresponding to (A) are re-colored by a specific Temporal-Color Code shown in the picture. (C) The intensity profiles of YFP and Chl autofluorescence in the white closed box shown in (A). A red asterisk indicates a spatial gap between the HR cell death lesion and *pVSP1* active domain. (D) Kymographs corresponding to the red box along the E–F axis in (B). (E) A magnified view of the yellow dashed box in (D). A spatial gap between the HR cell death lesion and *pVSP1* active domain is indicated by a red bar and a red arrow. (See also Supplementary Fig. S3; Supplementary Movie S3).

by two respective marker gene promoter activities in early *RPS2*-triggered immunity. Since the loss of SA accumulation by constitutive expression of *NahG* compromises not only *PR1* activation but also HR cell death, these local events are mainly considered to be under the control of SA accumulated around the infection foci (Delaney et al. 1994). The SA concentration gradient formed around the infection center appears to be rather steep not only in *RPS2*-conditioned ETI (Fig. 4E), but also in tobacco ETI against a viral pathogen (Enyedi et al. 1992),

indicating the presence of strict spatial regulation of the SA activation conserved in ETI. Considering the well-characterized mutually antagonistic relationship between SA and JA across multiple plant species (Spoel et al. 2007, Vlot et al. 2009), the JA active domain outside the SA active infection foci might contribute to limit overactivation of the SA pathway spatially around the infection site. This hypothesis should be further tested carefully by means of genetics and cell biology. In addition, the outer JA active domain may constitute another layer

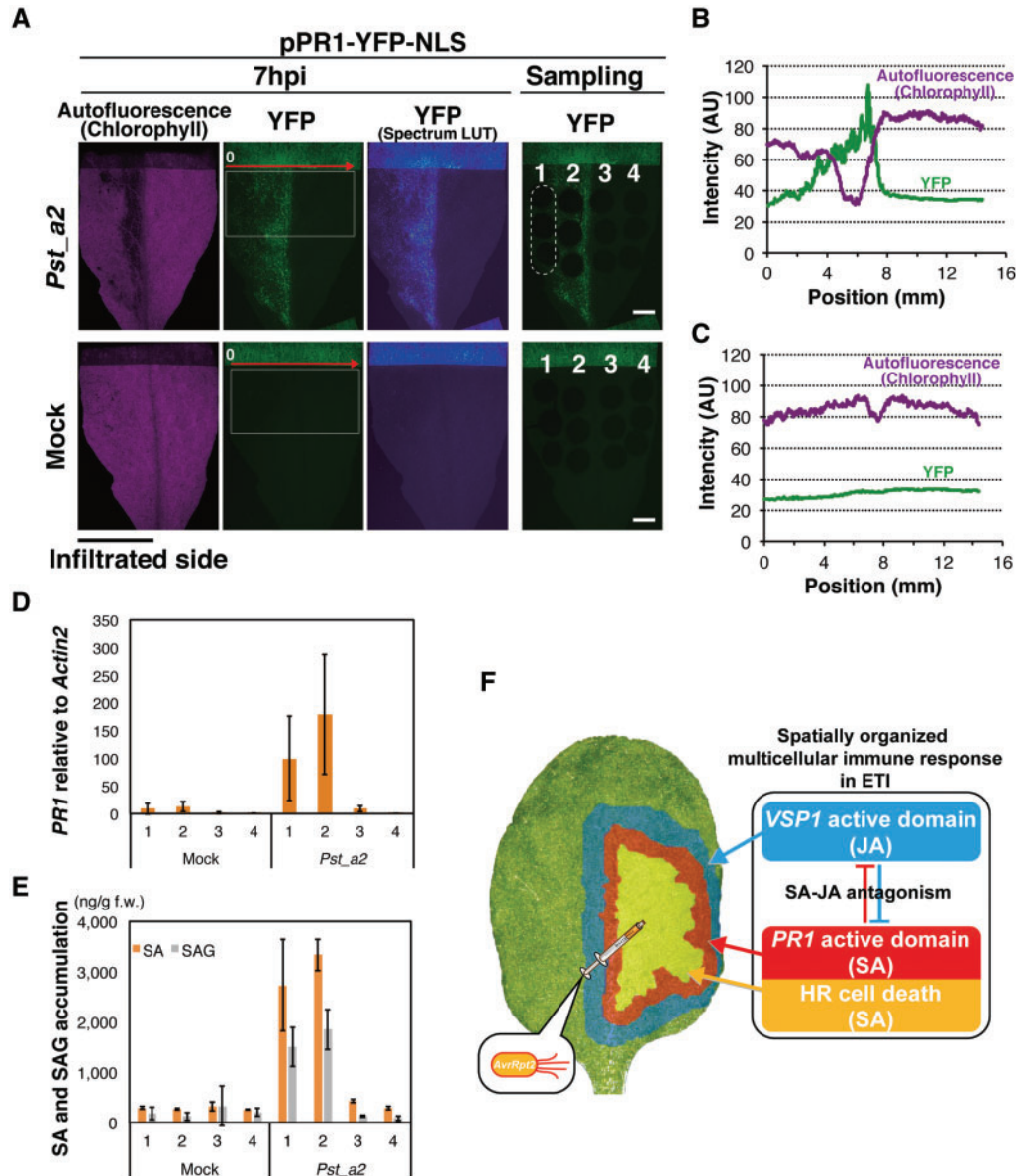


Fig. 4 Spatial regulation of SA accumulation leading to an organized multicellular response in ETI. (A) Site-specific sampling for SA and *PR1* analyses at 7 h.p.i. A nearly complete half of a pPR1-YFP-NLS leaf was fully infiltrated with *Pst_a2* (OD₆₀₀ = 0.2, upper) or 10 mM MgCl₂ (mock, lower). The leaf was divided into four areas along the mid-rib (numbered 1–4), and three leaf disks (2 mm in diameter) per area were sampled, as shown in a dashed ellipse for zone 1 in the upper right pictures. Representative sample pictures are shown. Scale bars = 2.5 mm. (B) Intensity profiles of YFP and autofluorescence in the white boxes in the *Pst_a2*-treated leaf in (A) along the red arrow. (C) Intensity profiles of YFP and autofluorescence in the white boxes in the mock-treated leaf in (A) along the red arrow. (D) The endogenous *PR1* expression levels in the four zones were measured by qRT-PCR. Eighteen leaf disks, corresponding to one zone, from six leaves were pooled as one sample. Bars represent means ± SD of three biological replicates. (E) The free SA and SA glycoside (SAG) levels in the four zones. Three disks from one zone from one leaf were pooled and analyzed. Bars represent means ± SD of three leaves. Experiments were repeated twice with similar results. (F) A schematic summary of an organized concentric pattern of the inner SA and the outer JA active domains which appeared around the infection site of *Pst_a2*. (See also Supplementary Fig. S4).

of ETI outside the SA domain, since the JA pathway also contributes positively to ETI conditioned by *RPS2* (Tsuda et al. 2009, Liu et al. 2016). The SA active cells undergoing programmed cell death during HR could be a target of necrotrophic pathogens (Spoel et al. 2007, Liu et al. 2016). The JA active domain surrounding the central SA active domain may have a function to protect living plant cells around the necrotic HR lesion from such secondary infections of necrotrophs. Taken

together, this virtually concentric pattern of the inner SA and the outer JA active domains found in this study may constitute a field of cells expressing orchestrated and comprehensive defense responses around the infection site during ETI. The biological relevance of this spatially organized ETI active field formation around the infection site now needs to be dissected further in detail. Another open question is whether or not this SA–JA concentric pattern in the ETI active field, which emerged

around the infection site, is only specific to ETI conditioned by RPS2. It will be worth investigating other pathosystems including non-ETI-causing pathogens using these promoter–reporters.

Our imaging-based analysis shed light on the importance of spatial aspects in understanding the complex plant immune signaling. Liu et al. (2016) demonstrated that the early activation of the JA pathway requires SA through SA receptors, instead of the conventional JA pathway (Liu et al. 2016). However, our data indicated that *VSP1* activation precedes *PR1*. More detailed genetic, biochemical and imaging-based studies, including other ETI responses, are required to understand how the concentric SA and JA active domains are formed from the spatiotemporal point of view. In addition, our data suggested, for the first time, that an apparent transient *PR1* activation is achieved through highly variable *PR1* activation in individual cells. Further detailed analysis at the single-cell level would answer the question of how plant tissues organize variable individual cellular activities into such collective behavior to form the concentric SA and JA active domains around the infection center. The complex activation of *PR1* and *VSP1* in the uninfected side of the leaves could also be further studied by single-cell level analyses. Thus, our imaging-based study stimulates further research to explore plant immunity spatiotemporally.

Materials and Methods

Plant materials and growth conditions

The *A. thaliana* wild type used in this study was Col-0. Water-soaked seeds were sown on soil and grown in a growth room at 23 °C under continuous white light (20–50 mmol m⁻² s⁻¹). Two- to three-week-old plants were used for all the experiments in this study.

Construction of transgenic promoter reporter plants

The 4.5 kb promoter of the *PR1* gene and the 3.0 kb promoter of the *VSP1* gene, both of which covered the previously analyzed respective regulatory sequences, were amplified from the genomic DNA (Col-0) by PCR and cloned into the pENTR/D-TOPO vector (Invitrogen) (Lebel et al. 1998, Utsugi et al. 1998). Primers used for the cloning are listed in [Supplementary Table S1](#). The promoter regions were recombined with the aid of Gateway technology into the binary pBGYN vector (Kubo et al. 2005). The resulting pBGYN-pPR1-YFP-NLS and pBGYN-pVSP1-YFP-NLS vectors were introduced into *Agrobacterium tumefaciens* GV3101::pMP90 and then into *A. thaliana* Col-0 wild-type plants using the floral dip method (Clough and Bent 1998). The characteristic spatial patterns of *pPR1* and *pVSP1* activities upon *Pst_a2* infiltration, shown in this study, were confirmed in multiple T₁ plants. Three (pPR1-YFP-NLS) and two (pVSP1-YFP-NLS) homozygous lines were selected by segregation analysis in the following generations. A representative homozygous line was selected for each for further detailed analyses.

Pseudomonas inoculation

Pst_a2 was previously described (Aarts et al. 1998). The bacterial cells were harvested and resuspended in 10 mM MgCl₂ to appropriate optical densities measured by a DU640 spectrophotometer (Beckman). The bacterial suspensions were infiltrated by hand into leaves using a 1 ml needleless syringe (Terumo).

Time-lapse imaging

Time-lapse imaging was performed using an M205FA automated stereomicroscope controlled by AF6000 software (Leica Microsystems). A DFC365FX camera (Leica Microsystems) was used in 12-bit mode. Chl autofluorescence and YFP were detected through TXR and YFP filters, respectively (both Leica

Microsystems). The TXR filter enabled us to reduce almost fully non-specific autofluorescence from dead cells. Bright field, YFP and TXR pictures were taken every 3 min and the intervals were programmed to expose the plant specimen to the light from the light-emitting diode (LED). Data analyses were performed with AF6000 (intensity plots) and Fiji (intensity plots, kymographs and Temporal-Color Code, ver. 2.0.0-rc-12/1.49g, build. 2352160d02) software.

Confocal microscopy

Confocal images were taken using a confocal microscope FV1200 equipped with UPLSAPO10 ×2, NA: 0.40 (Olympus). Z-projected pictures were generated by the FV10-ASW (Olympus). Chl autofluorescence was captured through propidium iodide (PI) and red fluorescent protein (RFP) pre-setting. Enhanced green fluorescent protein (eGFP) pre-setting was used for YFP imaging.

Gene expression analysis

Total RNA was isolated using an RNeasy Plant Mini Kit (Qiagen). For the first-strand cDNA synthesis using Superscript III, 100 ng (leaf disk samples) or 150 ng (whole-leaf samples) of total RNA were used. Quantitative reverse transcription–PCR (qRT–PCR) analysis was performed with a LightCycler TaqMan Master (Roche Applied Science) on a LightCycler 480 instrument II (Roche Applied Science). Relative mRNA levels were determined using *ACT2* as a reference gene. *PR1*, *VSP1* and *ACT2* expression was measured using UPLs #135, #91 and #30, respectively, with the primers listed in [Supplementary Table S1](#).

Quantification of hormone levels

The quantification of SA and SA glucoside (SAG) was performed as described previously with a minor modification (Seo et al. 1995). Briefly, three leaf disks were frozen and ground using liquid nitrogen. SA and SAG were extracted with 90% methanol, and SAG was converted to SA by β-glucosidase treatment. After separation by HPLC (Shimadzu) with an ODS column (μ-Bondasphere C18, 150 mm × ID3.9 mm, 5 μm, 100Å; Waters), SA levels were determined using a fluorescence detector (RF-20A; Shimadzu) with an excitation wavelength of 313 nm and an emission wavelength of 405 nm.

JA quantification was performed as described previously (Kojima et al. 2009, Shinozaki et al. 2015). Briefly, leaf samples (one leaf per sample) were frozen and ground using liquid nitrogen, and freeze dried. JA was extracted and determined using an ultra-HPLC-Q-Exactive™ system (Thermo Scientific) using an ODS column (AQUITY UPLC BEH C18, 1.7 μm, 2.1 × 100 mm; Waters) as described (Shinozaki et al. 2015).

Supplementary Data

Supplementary data are available at PCP online.

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Disclosures

The authors have no conflicts of interest to declare.

References

- Aarts, N., Metz, M., Holub, E., Staskawicz, B.J., Daniels, M.J. and Parker, J.E. (1998) Different requirements for *EDS1* and *NDR1* by disease resistance genes define at least two R gene-mediated signaling pathways in *Arabidopsis*. *Proc. Natl. Acad. Sci. USA* 95: 10306–10311.
- Bennett, M., Gallagher, M., Fagg, J., Bestwick, C., Paul, T., Beale, M., et al. (1996) The hypersensitive reaction, membrane damage and accumulation of autofluorescent phenolics in lettuce cells challenged by *Bremia lactucae*. *Plant J.* 9: 851–865.
- Berger, S., Bell, E., Sadka, A. and Mullet, J.E. (1995) *Arabidopsis thaliana* *Atvsp* is homologous to soybean *VspA* and *VspB*, genes encoding vegetative storage protein acid phosphatases, and is regulated similarly by methyl jasmonate, wounding, sugars, light and phosphate. *Plant Mol. Biol.* 27: 933–942.
- Clough, S.J. and Bent, A.F. (1998) Floral dip: a simplified method for *Agrobacterium*-mediated transformation of *Arabidopsis thaliana*. *Plant J.* 16: 735–743.
- Delaney, T.P., Uknes, S., Vernooij, B., Friedrich, L., Weymann, K., Negrotto, D., et al. (1994) A central role of salicylic acid in plant disease resistance. *Science* 266: 1247–1250.
- Dodds, P.N. and Rathjen, J.P. (2010) Plant immunity: towards an integrated view of plant–pathogen interactions. *Nat. Rev. Genet.* 11: 539–548.
- Dorey, S., Baillieul, F. and Pierrel, M.A. (1997) Spatial and temporal induction of cell death, defense genes, and accumulation of salicylic acid in tobacco leaves reacting hypersensitively to a fungal glycoprotein elicitor. *Mol. Plant* 10: 646–655.
- Enyed, A.J., Yalpani, N., Silverman, P. and Raskin, I. (1992) Localization, conjugation, and function of salicylic acid in tobacco during the hypersensitive reaction to tobacco mosaic virus. *Proc. Natl. Acad. Sci. USA* 89: 2480–2484.
- Fu, Z.Q., Yan, S., Saleh, A., Wang, W., Ruble, J., Oka, N., et al. (2012) NPR3 and NPR4 are receptors for the immune signal salicylic acid in plants. *Nature* 486: 228–232.
- Glazebrook, J. (2005) Contrasting mechanisms of defense against biotrophic and necrotrophic pathogens. *Annu. Rev. Phytopathol.* 43: 205–227.
- Guadagno, C.R., Ewers, B.E., Speckman, H.N., Aston, T.L., Huhn, B.J., DeVore, S.B., et al. (2017) Dead or alive? Using membrane failure and chlorophyll fluorescence to predict mortality from drought. *Plant Physiol.* 175: 223–234.
- Guerineau, F., Benjdia, M. and Zhou, D.X. (2003) A jasmonate-responsive element within the *A. thaliana* *vsp1* promoter. *J. Exp. Bot.* 54: 1153–1162.
- Kazan, K. and Manners, J.M. (2013) MYC2: the master in action. *Mol. Plant* 6: 686–703.
- Kenton, P., Mur, L.A.J., Atzorn, R., Wasternack, C. and Draper, J. (1999) (–)-Jasmonic acid accumulation in tobacco hypersensitive response lesions. *Mol. Plant Microbe Interact.* 12: 74–78.
- Kojima, M., Kamada-Nobusada, T., Komatsu, H., Takei, K., Kuroha, T., Mizutani, M., et al. (2009) Highly sensitive and high-throughput analysis of plant hormones using MS-probe modification and liquid chromatography–tandem mass spectrometry: an application for hormone profiling in *Oryza sativa*. *Plant Cell Physiol.* 50: 1201–1214.
- Kubo, M., Udagawa, M., Nishikubo, N., Horiguchi, G., Yamaguchi, M., Ito, J., et al. (2005) Transcription switches for protoxylem and metaxylem vessel formation. *Genes Dev.* 19: 1855–1860.
- Lebel, E., Heifetz, P., Thorne, L., Uknes, S., Ryals, J. and Ward, E. (1998) Functional analysis of regulatory sequences controlling PR-1 gene expression in *Arabidopsis*. *Plant J.* 16: 223–233.
- Liu, L., Sonbol, F.-M., Huot, B., Gu, Y., Withers, J., Mwimba, M., et al. (2016) Salicylic acid receptors activate jasmonic acid signalling through a non-canonical pathway to promote effector-triggered immunity. *Nat. Commun.* 7: 13099.
- Mine, A., Sato, M. and Tsuda, K. (2014) Toward a systems understanding of plant–microbe interactions. *Front. Plant Sci.* 5: 423.
- Murray, S.L., Thomson, C., Chini, A., Read, N.D. and Loake, G.J. (2002) Characterization of a novel, defense-related *Arabidopsis* mutant, *cirt*, isolated by luciferase imaging. *Mol. Plant Microbe Interact.* 15: 557–566.
- Ohshima, M., Itoh, H., Matsuoka, M., Murakami, T. and Ohashi, Y. (1990) Analysis of stress-induced or salicylic acid-induced expression of the pathogenesis-related 1a protein gene in transgenic tobacco. *Plant Cell* 2: 95–106.
- Schmelzer, E., Kruger-Lebus, S. and Hahlbrock, K. (1989) Temporal and spatial patterns of gene expression around sites of attempted fungal infection in parsley leaves. *Plant Cell* 1: 993–1001.
- Seo, S., Okamoto, M., Seto, H., Ishizuka, K., Sano, H. and Ohashi, Y. (1995) Tobacco MAP kinase: a possible mediator in wound signal transduction pathways. *Science* 270: 1988–1992.
- Shinozaki, Y., Hao, S., Kojima, M., Sakakibara, H., Ozeki-Iida, Y., Zheng, Y., et al. (2015) Ethylene suppresses tomato (*Solanum lycopersicum*) fruit set through modification of gibberellin metabolism. *Plant J.* 83: 237–251.
- Spoel, S.H., Johnson, J.S. and Dong, X. (2007) Regulation of tradeoffs between plant defenses against pathogens with different lifestyles. *Proc. Natl. Acad. Sci. USA* 104: 18842–18847.
- Spoel, S.H., Koornneef, A., Claessens, S.M., Korzelius, J.P., Van Pelt, J.A., Mueller, M.J., et al. (2003) NPR1 modulates cross-talk between salicylate- and jasmonate-dependent defense pathways through a novel function in the cytosol. *Plant Cell* 15: 760–770.
- Tsuda, K. and Somssich, I.E. (2015) Transcriptional networks in plant immunity. *New Phytol.* 206: 932–947.
- Tsuda, K., Mine, A., Bethke, G., Igarashi, D., Botanga, C.J., Tsuda, Y., et al. (2013) Dual regulation of gene expression mediated by extended MAPK activation and salicylic acid contributes to robust innate immunity in *Arabidopsis thaliana*. *PLoS Genet.* 9: e10004015.
- Tsuda, K., Sato, M., Stoddard, T., Glazebrook, J. and Katagiri, F. (2009) Network properties of robust immunity in plants. *PLoS Genet.* 5: e1000772.
- Utsugi, S., Sakamoto, W., Murata, M. and Motoyoshi, F. (1998) *Arabidopsis thaliana* vegetative storage protein (VSP) genes: gene organization and tissue-specific expression. *Plant Mol. Biol.* 38: 565–576.
- Vlot, A.C., Dempsey, D.A. and Klessig, D.F. (2009) Salicylic acid, a multifaceted hormone to combat disease. *Annu. Rev. Phytopathol.* 47: 177–206.
- Wolpert, L. (1969) Positional information and the spatial pattern of cellular differentiation. *J. Theor. Biol.* 25: 1–47.
- Yan, S. and Dong, X. (2014) Perception of the plant immune signal salicylic acid. *Curr. Opin. Plant Biol.* 20: 64–68.
- Yu, G.L., Katagiri, F. and Ausubel, F.M. (1993) *Arabidopsis* mutations at the *RPS2* locus result in loss of resistance to *Pseudomonas syringae* strains expressing the avirulence gene *avrRpt2*. *Mol. Plant Microbe Interact.* 6: 434–443.
- Zeier, J., Pink, B., Mueller, M.J. and Berger, S. (2004) Light conditions influence specific defence responses in incompatible plant–pathogen interactions: uncoupling systemic resistance from salicylic acid and PR-1 accumulation. *Planta* 219: 673–683.