Post-translational regulation of plant immunity
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Plants have evolved multi-layered molecular defense strategies to protect against pathogens. Plant immune signaling largely relies on post-translational modifications (PTMs) to induce rapid alterations of signaling pathways to achieve a response that is appropriate to the type of pathogen and infection pressure. In host cells, dynamic PTMs have emerged as powerful regulatory mechanisms that cells use to adjust their immune response. PTM is also a virulence strategy used by pathogens to subvert host immunity through the activities of effector proteins secreted into the host cell. Recent studies focusing on deciphering post-translational mechanisms underlying plant immunity have offered an in-depth view of how PTMs facilitate efficient immune responses and have provided a more dynamic and holistic view of plant immunity.

Introduction

Plants are in constant association with microbial communities, and have evolved a multi-layered molecular defense strategy to protect themselves against pathogens. In the absence of specialized immune cells, plants rely on rapid alterations of signaling pathways within individual cells to achieve an appropriate defense response while balancing the trade-off with normal growth and metabolism. Much of our knowledge about the regulation of plant immune responses has been obtained using genetic and genomic approaches for identification of genes and gene expression patterns underlying defense pathways. However, genetic approaches are limited to examining the effects of losing or gaining gene functions, rather than dynamic regulation of cellular processes. Consequently, our current understanding of plant immune networks is far from complete. In recent years, there has been a significant shift from studying the underlying transcriptional networks to studying protein post-translational modification (PTM), which is a versatile regulatory process that rapidly alters the functional diversity of the proteome [1]. Current evidence shows that PTMs are critical for the rapid reprogramming of cells for defense signaling and for attenuating the response to achieve cellular homeostasis in all layers of the plant immune responses.

In this review, we seek to highlight discoveries that have demonstrated the power and versatility of PTMs as regulatory mechanisms for host cells to rapidly respond to pathogens and for pathogens to deploy effectors as a part of their virulence strategy.

Phosphorylation dynamics during pathogen perception and pattern-triggered immunity

At the front line of molecular defense strategies, plants deploy a molecular surveillance system through the cell surface-anchored pattern-recognition receptors (PRRs) to detect microbe-associated and damage-associated molecular patterns (MAMPs and DAMPs) and initiate pattern-triggered immunity (PTI) (Figure 1) [2]. PTI is effective against a broad range of microbes and is characterized by rapid calcium influx, production of reactive oxygen species (ROS), mitogen activated protein kinase (MAPK) phosphorylation cascades, callose deposition, and defense gene expression [3]. The activation of PTI also leads to plant growth inhibition, highlighting the trade-off between growth and defense physiology. Therefore, understanding how this immune response is dynamically regulated by PTM to achieve rapid and transient induction has been a major focus in deciphering the mechanisms of PTI.

Activation of the PRRs, FLAGELLIN-SENSITIVE 2 (FLS2) and EF-TU RECEPTOR (EFR), by peptides derived from bacterial flagellin (flg22) and elongation factor Tu (elf18), stimulates the recruitment of the leucine-rich receptor-like kinase (LRR-RLK) BR1-ASSOCIATED RECEPTOR KINASE (BAK1) and the receptor-like cytoplasmic kinase (RLCK) BOTRYTIS-INDUCED KINASE1 (BIK1). These plasma membrane (PM)-associated kinases are in turn activated by phosphorylation to initiate defense signal transduction [3,4]. BAK1 and BIK1 are shared by many PRR signaling pathways, including the growth hormone brassinosteroid (BR) signal mediated by the receptor BRASSINOSTEROID INSENSITIVE 1 (BR1) [5]. Recent studies using targeted phospho-proteomics have revealed differential phosphorylation patterns for BAK1 in complex with distinct RLKs (i.e., FLS2, EFR, BIK1 and BR1). For example, phosphorylation of BAK1 by BIK1 at four amino
acids in the kinase domain and two in the activation loop is inhibited through phosphorylation mediated by FLS2 at Ser-286 [6**]. A similar inhibitory effect was observed for EFR through phosphorylation of Thr-455 in the BAK1 activation domain [6**]. The opposing activities of FLS2 and EFR against BIK1-mediated phosphorylation suggest that once BAK1 and BIK1 have become phosphorylated in response to MAMP perception, the PRRs can attenuate this immune response through inhibition of BAK1 phosphorylation mediated by BIK1.

Besides the opposing activities between BIK1 and FLS2/EFR on BAK1, phosphatases, such as SERINE/THREONINE PROTEIN PHOSPHATASE 2A (PP2A), can also counteract the kinase activity to suppress PTI [7]. Upon flg22 treatment, PP2A activity has to be reduced to allow induction of BAK1 activity and PTI signaling [7]. Additionally, the PM-localized PP2C38 interacts with FLS2-BIK1 as well as EFR-BIK1 and dephosphorylates BIK1 to inhibit PAMP-induced ROS production and stomatal closure [8**]. Co-immunoprecipitation (Co-IP) experiments demonstrated that PP2C38 and BIK1 directly interact with each other. A combined approach using genetics and phospho-proteomic analysis indicated that MAMP treatment induces BIK1-dependent phosphorylation of PP2C38 at Ser-77, which significantly diminishes the PP2C38 activity and its interaction with the receptor complexes [8**]. This interplay between immune regulatory kinases and phosphatases is a major mechanism by which plant cells achieve homeostasis during infection.

Phosphorylation dynamics are essential for regulation of immune responses; therefore, they are also prime targets for the activity of bacterial type III effector proteins (TTEs). Pseudomonas syringae pv tomato DC3000 (Pst DC3000) delivers a protein tyrosine phosphatase (PTP), HopAO1, into host cells as part of its virulence...
strategy. When HopAO1 is ectopically expressed in *Arabidopsis*, it inhibits elf18- and flg22-induced ROS burst, MAPK activation, and resistance to *Pst* DC3000 [9]. HopAO1 interacts with the kinase domains of FLS2 and EFR in yeast and plant cells, and in vitro kinases assays demonstrated that the presence of HopAO1 results in a significant decrease in elf18-induced Tyr phosphorylation on EFR [9]. The ability of *Pst* DC3000 to colonize host tissue is compromised in strains harboring a HopAO1 deletion, and this defect is absent when inoculated into *Arabidopsis* fls2/efr double mutants, confirming that a major virulence function of HopAO1 is the targeted dephosphorylation of PRRs to inhibit MAMP-induced immune responses [9].

**Ubiquitination and N-glycosylation contribute to PTI through degradation and cellular targeting of PRRs**

PRR signaling dynamics are fine-tuned by ubiquitination, which facilitates their endocytosis and degradation by the 26S proteasome [10,11]. Yeast two-hybrid (Y2H) and Co-IP experiments using the BAK1 kinase domain as a bait demonstrated that flg22 treatment stimulates interaction between BAK1 and the plant U box (PUB) E3 ubiquitin ligases PUB12 and PUB13, and that the BAK1-PUB12/13 complex associates with FLS2 within 30 s of flg22 treatment [12]. Kinase assays using protoplasts expressing BAK1 or a kinase-inactive mutant indicated that BAK1 phosphorylates the C-terminal Armadillo (ARM) repeat domain of PUB13 in response to flg22 treatment, and chemical inhibition of kinase activity confirmed that BAK1-dependent phosphorylation is required for flg22-induced FLS2-PUB12/PUB13 interaction [12,13]. *In vitro* ubiquitination assays confirmed that PUB12 and PUB13 polyubiquitinate the cytosolic domain of FLS2, but not BAK1 or BIK1 [12]. Three additional, functionally redundant U-box E3 ubiquitin ligases, PUB22/23/24 have also been shown to negatively regulate PTI responses [14]. Upon flg22 perception, PUB22 ubiquitinates Exo70B2, a member of a vesicle tethering complex, and targets it for degradation; Exo70B2 is required for both early and late PTI responses [15,16]. Taken together, studies of ubiquitin-mediated regulation of PRRs indicate that their internalization and degradation likely serves to desensitize cells to continual MAMP stimulus, as well as to reset cells for new rounds of signaling [17].

Asparagine-linked protein glycosylation (N-glycosylation) contributes to protein folding and stability, secretion, and interactions with ligands and other proteins [18]. Treatment of *Arabidopsis* protoplasts with tunicamycin, which blocks N-glycosylation, and subsequent biochemical analyses showed that the extracellular LRR domains of both FLS2 and EFR are extensively N-glycosylated and that these modifications are required for their maturation and transport to the PM [19]. Site-directed mutagenesis has also been used to generate point mutations in EFR that disrupt individual N-X-(S/T) glycosylation motifs and affect discrete N-glycosylation events. When transiently expressed in *Nicotiana benthamiana*, which is deficient in elf18 perception, the point mutant of EFR, Asn-143-Gln, fails to elicit MAMP-induced ROS production [19], indicating that individual N-glycosylation sites also play a role in PRR stability and ligand recognition.

**Downstream of the PRRs—MAPK cascades and production of reactive oxygen species**

Upon pattern recognition and activation of PRR-RLK complexes, immune signaling is propagated by MAPK phosphorylation cascades leading to transcriptional reprogramming for defense [2]. Recent studies of the FLS2 signaling complex have revealed a connection between PRR/RLK signals and the MAPK phosphorylation cascade, which has been a missing link in PTI signaling against bacterial pathogens. MITOGEN-ACTIVATED PROTEIN KINASE KINASE KINASE 7 (MKKK7) was first identified in a large-scale proteomics screen for differential protein phosphorylation upon flg22 treatment [20]. MKKK7 interacts directly with FLS2, and quantitative phospho-proteomic analyses indicated that upon flg22 treatment, MKKK7 is phosphorylated at two different serine residues in a temporally distinct manner [21**]. MKKK7 appears to be a negative regulator of PTI because mkkk7 mutants enhanced MPK6 phosphorylation, increased expression of flg22-induced WRKY29 and FRK1, and enhanced resistance to *Pst* DC3000 [21**]. The upstream kinases that act on MKKK7 have yet to be firmly established.

Phosphatases that regulate the MAPK cascade include PROTEIN TYROSINE PHOSPHATASE 1 (PTP1) and the dual-specificity phosphatase (DSP) MKP1, which target MPK3 and MPK6, and down-regulate defense responses against *Pst* DC3000 [22–24]. Members of the PROTEIN PHOSPHATASE 2C (PP2C) family, AP2C1 and AP2C2, were recently shown to be negative regulators of PTI against *P. syringae*. The *ap2c1* mutants exhibited enhanced activation of MPK3, 4, and 6 in response to flg22 and elf18 treatment, as well as to *Pst* DC3000 infection, resulting in upregulation of genes encoding SA biosynthetic enzymes, WRKY TFs, MAPKK/MAPK kinases, and ultimately enhanced callose deposition and bacterial resistance [25].

Another hallmark of PTI is the production of reactive oxygen species (ROS), which have antimicrobial activities and contribute to defense signaling, cell-wall reinforcement, and stomatal closure to limit pathogen entry into host tissues [26,27]. The respiratory burst oxidase homolog (RBOH) proteins, which are PM-localized NADPH oxidases, are required for PTI-triggered ROS production and are activated through phosphorylation by calcium-dependent protein kinases (CDPKs) and the RLCK BIK1 (Figure 1) [26,28–30]. ROS signaling is
linked to metabolic pathways important for defense against pathogens. GLUCOSE-6-PHOSPHATE DEHYDROGENASE (G6PD), a key enzyme of the oxidative pentose phosphate pathway, is regulated by cellular redox status and phosphorylation. G6PD generates NADPH and metabolic intermediates for numerous biosynthetic pathways and exhibits pathogen-inducible activity that is required for the ROS burst [31,32]. The Arabidopsis GLYCOGEN SYNTHASE KINASE3 (GSK3)/Shaggy-like kinase ASKα, well-known for its role in regulating abiotic stress and brassinosteroid signaling responses, also promotes PTI [31,33**]. Treatment of seedlings with multiple MAMPs including fliG22, chitin, and the DAMP PEPTIDE 1 (PEP1), rapidly induces ASKα protein levels and enzymatic activity. Plants over-expressing ASKα display increased ROS production and transcription of PTI marker genes, while these responses are attenuated in aska mutants [33**]. In response to abiotic stress, ASKα phosphorylates G6PD6 at Thr-467 [31], and expression of phospho-mimetic (T467E)/phospho-null (T467A) G6PD6 variants in Arabidopsis revealed that phosphorylation at this residue is also critical for enhanced ROS production. Furthermore, infection by Pst DC3000 stimulates ASKα expression levels and kinase activity, and both aska and gpd6 result in increased susceptibility to the bacterial infection while expression of wild type G6PD6 or T467E restores resistance [33**]. The results from this work suggest that upon MAMP treatment, ASKα phosphorylates G6PD6 at Thr-467 leading to increased ROS production and enhanced PTI [31,33**].

ROS can also influence growth-defense balance through crosstalk among hormones such as indole-3-acetic acid (IAA) for growth and salicylic acid (SA) and jasmonic acid (JA) for defense against biotrophic and necrotrophic pathogens, respectively. The H2O2 scavenging enzyme CATALASE 2 (CAT2) was recently shown to participate in SA-mediated repression of IAA biosynthesis during defense responses [34,35**]. SA binds to CAT2 and inhibits its activity resulting in H2O2 accumulation [35**]. The increase in H2O2 promotes sulfenylation of Tryptophan (Trp) Synthetase β subunit 1 (TSB1) to inhibit its activity leading to reduction of Trp levels and ultimately decreased auxin production [35**]. This PTM of TSB1 likely occurs on Cys-308 as indicated by inhibited enzymatic activity and the insensitivity of Cys-308-Ser to H2O2-mediated sulfenylation [35**]. These conclusions were supported by the finding that over-expression of TSB1 could rescue the reduced Trp and IAA concentrations in the cat2-1 mutant [35**]. CAT2 also interacts with and promotes the activity of ACYL CO-ENZYM A OXIDASE (ACX2 and ACX3), peroxisomal enzymes in the JA biosynthetic pathway. Treatment with SA, or induction of SA biosynthesis in response to biotrophic pathogens significantly inhibits the interaction among these enzymes leading to reduction in JA biosynthesis [35**]. Since some pathogens have the ability to stimulate IAA biosynthesis in plant tissues to facilitate infection [34] and/or take advantage of inhibitory SA/JA crosstalk in their favor, this work has revealed a mechanism by which host cells can counter these virulence strategies through SA-mediated inhibition of IAA and JA biosynthesis during defense signaling.

Post-translational dynamics during effector-triggered immunity and systemic acquired resistance

PTI can be overcome by pathogens through delivery of TTSS into host cells. The virulence function of effectors is often achieved through PTM of host proteins to manipulate immune signaling cascades [36]. In response, plants have evolved resistance (R) proteins, which are intracellular nucleotide-binding leucine-rich repeat (NB-LRR) receptors that either directly or indirectly detect the activity of effector proteins to initiate effector-triggered immunity (ETI). ETI commonly leads to calcium and ROS accumulation, massive transcriptional reprogramming, and induction of programmed cell death (PCD) at the site of infection that limits the spread of the pathogen [37–40]. ETI also triggers the biosynthesis of the defense hormone SA and expression of anti-microbial pathogenesis related (PR) proteins in infected (local) and systemic tissue leading to establishment of systemic acquired resistance (SAR), which protects against additional infection [41].

PTMs at the host-pathogen interface

The Arabidopsis RPM-1 INTERACTING PROTEIN 4 (RIN4) is a multi-functional immune response regulator at the host-pathogen interface that is targeted by at least five P. syringae effectors and guarded by the host NB-LRRs RPM1 and RPS2 [42–45]. RPM1 could detect phosphorylation of RIN4-Thr166 by the effector (AvrB and AvrRpm1)-mediated expression and activation of a host RLCK, RPK [44,46]. In this case, the perturbation caused by the effectors are detected by the cognate host NB-LRRs to trigger ETI. However, effectors normally evolved to help the pathogens. For example, a bacterial cysteine protease, AvrPphB, contributes to the dynamics by cleaving RIPK, thereby suppressing RPM1-mediated ETI, resulting in increased susceptibility to Pst DC3000 expressing AvrB [47*].

The host RIN4 also serves as a regulatory link between ETI and PTI. Upon fliG22 perception, RIN4 is rapidly phosphorylated at Ser-141 leading to enhanced PTI [48]. This phosphorylation event is upstream of and independent from phosphorylation of RIN4 at Thr-166 that activates ETI in response to AvrB/AvrRpm1 and does not affect the cleavage of RIN4 by AvrRpt2 [48]. Phosphorylation of RIN4-Thr-166, which represses PTI signaling, is reduced in response to fliG22 treatment but enhanced in the presence of AvrB offering insight into
the mechanism by which *P. syringae* subverts the host’s initial defense responses [49**]. Additionally, triple phospho-mimic RIN4 mutants (T21D, S160D, or T166D) exhibit enhanced interaction with, and activation of the plasma membrane H+-ATPase 1 (AHA1), and consequently wider stomatal apertures and enhanced colonization of spray inoculated *P. syringae* [49**]. Although the evidence demonstrates that RIN4 pSer-141 requires interaction with FLS2, the kinase responsible for this modification remains elusive.

In addition to targeting upstream signaling components, such as RIN4, some effectors have evolved to help pathogens evade immune responses through PTM of downstream components, such as TFs. Co-IP experiments demonstrated that PopP2 from *Ralstonia solanacearum* acetylates WRKY41,70, and 33, which are positive regulators of defense gene expression, and reduces their ability to bind to gene promoters resulting in suppression of host immune responses [50**,51**]. To counteract this pathogen virulence strategy the NB-LRR RESISTANCE TO *RALSTONIA SOLANACEARUM* 1 (RRS1) is activated directly through effector-mediated PTMs of its extra C-terminal WRKY DNA binding domain [50**,51**,52]. Biochemical and proteomics analyses revealed that PopP2 also acetylates RRS1 on four lysine residues in its WRKY domain with Lys-1221 being a key residue for this modification [50**,51**]. The WRKY

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**Figure 2**

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NPRs are regulators of two opposing hormone-mediated immune responses. NPR1 normally exists as a high molecular weight oligomer in the cytoplasm. In response to salicylic acid (SA)-induced changes in cellular redox, NPR1 monomers are released into the nucleus. At resting state, NPR1, which is phosphorylated (red circles) at Ser55/Ser59, interacts with WRKY, a transcriptional repressor of PR genes. Sumoylated (light blue oval) NPR1, which is phosphorylated at Ser11/Ser15 but dephosphorylated at Ser55/Ser59, interacts with the transcriptional activator TGA3 to promote PR gene expression and establishment of systemic acquired resistance (SAR). NPR3 and NPR4 mediate SA-dependent degradation of NPR1 by the proteasome. NPR3 and NPR4 also promote the SA-enhanced degradation of JAZ transcriptional repressors to induce JA gene expression, resulting in activation of JA synthesis and signaling during early effector-triggered immune (ETI) responses.
domain of RRS1-R from *Arabidopsis* accessions Ws-2 and Nd-1 has an extended C-terminus that is hypothesized to interact with negative regulatory *cis*-elements in the promoters of genes that initiate ETI [51**]. Therefore, PopP2-dependent acetylation of RRS1 would attenuate its ability to bind to W-box motifs present in defense gene promoters and consequently trigger ETI [50**,51**]. Taken together, the results of these two studies reveal that acetylation of WRKY TFs is a virulence function of PopP2, and that the integrated WRKY domain of RRS1 has evolved as a decoy for recognition of the effector activity and induction of ETI.

**NPRs are regulators of two opposing SA-mediated immune responses**

During ETI, the plant defense hormones against biotrophic and necrotrophic pathogens, SA and JA respectively, both accumulate in the infected tissue [53]. Although many studies support an antagonistic SA-JA relationship that depends on the transcription co-factor NPR1 [54,55], there is also considerable evidence indicating that these hormones have a cooperative role during ETI [56,57]. A recent study has now shown that JA plays a positive role in establishment ETI through a mechanism that does not require NPR1 or the canonical JA receptor, CORONATINE INSENSITIVE 1 (COI1), but requires the SA receptors NPR3 and NPR4. NPR3 and NPR4 interact with JAZ transcriptional repressors in yeast and *in planta*, and promote JAZ1 degradation during early stages of ETI to activate expression of JA biosynthesis and signaling genes (Figure 2) [58**]. Given that NPR3/NPR4 are substrate adapters for an E3 ubiquitin ligase complex, the evidence suggests that ubiquitination is the PTM leading to JAZ degradation by the proteasome during early ETI responses.

In contrast to ETI, which is a signal-specific local response, SAR confers broad-spectrum resistance in uninfected systemic tissues. NPR1 is both a positive regulator of SAR and a negative regulator of ETI and associated PCD (Figure 2) [41]. In addition to SA-induced, redox-sensitive localization dynamics, the activity of NPR1 in the nucleus is tightly regulated by PTMs [59]. The discoveries of two opposing phosphorylation events (Ser55/Ser59 inhibits and Ser11/Ser15 activates NPR1) and their interplay with sumoylation, a PTM affecting NPR1 differential associations with TFs, TGAs and WRKYs, have revealed complex posttranslational regulation of NPR1-dependent transcription during immune responses [60**]. Sumoylation of NPR1, which is phosphorylated at Ser11/Ser15 but dephosphorylated at Ser55/Ser59, is also required for interaction with NPR3 and NPR4 leading to its degradation by the proteasome [60**,61]. Differences in the SA binding affinities for NPR3 (low) and NPR4 (high) and the opposing effects of SA binding to their interactions with NPR1 (*i.e.*, SA facilitates NPR3–NPR1 interaction but disrupts NPR4–NPR1 interaction) [61], allows NPR1 degradation to occur in cells with high SA concentration (infection site) to remove inhibition on ETI and PCD. NPR1 is stabilized in systemic tissue where SA concentration is too low for NPR3–NPR1 interaction but high enough to disrupt NPR4–NPR1 interaction [61]. Thus, post-translational control of the NPR1 protein level allows PCD in local tissue and establishment of SAR in systemic tissues [61].

**Conclusions**

Several PTMs stand out as major players of immune regulation; notably, phosphorylation, ubiquitination, and sumoylation. Our understanding of other modifications such as S-nitrosylation, acetylation, and sulfenylation, and their importance to plant immune responses and pathogen virulence mechanisms is also expanding. Although phosphorylation is a prominent posttranslational mechanism for propagating rapid and transient immune responses, identifying the kinases upstream of a known substrate remains a major challenge. The use of targeted, quantitative proteomics and the application of proximity-dependent affinity labeling of protein complexes *in vivo* show promise for the discovery of unknown kinases [62,63**,64**]. Additionally, advances in the sensitivity of proteomics instrumentation is rapidly increasing our ability to detect PTMs that occur in low stoichiometry, and therefore will open the door to understanding how these modifications contribute to immune responses. Another exciting avenue for future studies, which will be paramount to understanding the dynamic immune system of plants, will be to connect protein kinases to their counteracting protein phosphatases and understand both the quantitative and the physiological significance of these opposing modifications. Gaining an in-depth understanding of how PTMs facilitate efficient immune responses and link PTI, ETI, and SAR will provide a dynamic and holistic view of plant immunity, and lead to new strategies for improving plant performance in the face of pathogen challenge.

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**References and recommended reading**

Papers of particular interest, published within the period of review, have been highlighted as:

- of special interest
- of outstanding interest


Using quantitative phospho-proteomics, this study revealed that BAK1 is differentially phosphorylated when in complex with different PRR complexes indicating that differential phosphorylation may be the molecular mechanisms for discrete regulation of BAK1-dependent pathways.


This study focuses on the negative regulation of PTI through the activity of the protein phosphatase Pp2c38 on the cysteine kinase BIK1, a component of several PRR complexes. Pp2c38 was demonstrated to be a negative regulator of BIK1 activity and PTI-mediated immunity through attenuation of PAMP-induced BIK1 phosphorylation, which decreases phosphorylation of the NADPH oxidase RboHD by BIK1.


This work identifies MAPKK7 as an FLS2-interacting kinase, providing a link between PRR complexes and the MAPK phosphorylation cascade. Targeted proteomics indicated that MKK7 is differentially phosphorylated at specific serine residues, and functional analyses demonstrated that this kinase negatively regulates flagellin-triggered immune responses through suppression of MKP6 activity, ROS production, and defense gene expression.


This study identifies GSK3/ASKα kinase as a positive regulator of plant immunity. GSK/ASKα directly phosphorylates and stimulates the activity of G6PD1, a key enzyme of the NADPH-dependent phosphate pathway.
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providing a link between protein phosphorylation cascades during PTI to primary metabolic pathways.


This study demonstrates that the redox scavenging enzyme CAT2 mediates the inhibitory effect of the defense hormone SA on auxin and JA biosynthesis. The authors demonstrated that SA suppresses CAT2 causing increases in H$_2$O$_2$ and sulfonylation of the tryptophan synthase enzyme TS1. Inactivation of TS1 by sulfonylation reduces tryptophan levels and limits this chemical precursor of auxin synthesis. CAT2 was also shown to have a positive effect on JA biosynthesis by promoting interaction between the JA biosynthetic enzymes, AOX and AOX3. Therefore, SA repression of CAT2 inhibits JA accumulation.


This study reveals that the bacterial effector protein AvrPphB suppresses AvrB-mediated activation of RPM1, but not AvrRpm1-mediated activation, through cleavage of RIPK by AvrPphB. This indicates the presence of distinct mechanisms for recognition of AvrB and AvrRpm1 by the single R protein RPM1.


The P. syringae effector AvrB targets the host proteins RIN4 and RIPK. RIPK phosphorylates RIN4 at Thr-166, which is sensed by the NB-LRR RPM1 leading to activation of ETI. This work demonstrated that phosphomimetic RIN4 mutants result in reduced flagellin-triggered responses, and enhanced susceptibility to P. syringae due to inhibition of stomatal defense through enhanced plasma membrane H(+)-ATPase activity. The authors demonstrate that basal phosphorylation of RIN4 Thr-166 decreases upon flagellin treatment, but not in the presence of AvrB, suggesting that AvrB targets RIN4 to enhance pathogen entry through stomata.


This study, in combination with the work from Le Roux et al. (2015), offers significant insight into how NB-LRRs act as a surveillance system for the activity of a pathogen effectors. Using two RRS1 alleles, RRS1-S (Arabidopsis Col-0) and RRS1-R (ArabidopsisWs-2 and Nd-1), the authors demonstrate that the integrated WRKY domain present at the C terminus acts as a decoy for the bacterial effector PopP2, which normally acetylates WRKY TFs that are positive regulators of defense. PopP2 acetylates the RRS1-S WRKY domain and inhibits AvrRps4 recognition. PopP2-dependent RRS1 acetylation reduces its ability to bind defense gene promoters. The authors suggest that the extended C terminus of RRS1-R is essential for PopP2-dependent ETI.


This study, in combination with the work from Sarris et al. (2015), offers significant insight into how NB-LRRs act as a surveillance system for the activity of a pathogen effectors. Using an RRS1 allele that is present in ArabidopsisWs-2 and Nd-1, RRS1-R, the authors demonstrate that the integrated WRKY domain present at the C terminus acts as a decoy for the biochemical activity of the bacterial effector PopP2, which normally acetylates WRKY TFs that are positive regulators of defense. PopP2 acetylates conserved lysines, specifically Lys-1231 in the RRS1-R WRKY domain, which reduces its ability to bind defense gene promoters and triggers ETI.


In this study, the authors provide strong evidence that a plant defense hormone, JA, is a positive regulator of RPS2-mediated ETI. They demonstrate that during ETI, early induction of JA synthesis following SA accumulation is activated through the SA receptors NPR3 and NPR4.
instead of the JA receptor COI1. NPR3 and NPR4 promote degradation of JAZ proteins, transcriptional repressors of JA signaling. This discovery offers a mechanistic understanding of how plants can defend against a biotrophic pathogen without increasing susceptibility to necrotrophic pathogens.


This work demonstrates that NPR1 is sumoylated by SUMO3, and that this PTM serves to activate defense responses by switching NPR1’s association with WRKY70 (transcriptional repressor) to TGA3 (transcriptional activator). This study also refines the model of NPR1 phosphorylation dynamics by showing that NPR1 is phosphorylated at Ser55/Ser59 during resting state, and that SA induces dephosphorylation of Ser55/Ser59, which is a requirement for NPR1 activation by sumoylation and phosphorylation at Ser11/Ser15.


Proteomics approaches to quantitative identification of PTMs have rapidly advanced in recent years. This work describes methodology for determining biologically relevant kinase substrates using a combination of techniques including selective chemical inhibition, quantitative phosphoproteomics, and machine learning. This platform was applied to identify direct targets of Cdc28 and Snf1 kinases in S. cerevisiae; however, the methods are broadly applicable and can be applied in other species to facilitate global phosphorylation network profiling.


Proximity-dependent labeling of interacting proteins using a promiscuous biotin ligase fused to a bait protein is a powerful technique that has the capability of identifying members of protein complexes, as well as rapid transient interactors such as kinases or phosphatases. This review describes the widely used BioID platform and its applications to different target protein types, and compares/contrasts this method to other proximity-dependent labeling approaches.