

Specific Threonine Phosphorylation of a Host Target by Two Unrelated Type III Effectors Activates a Host Innate Immune Receptor in Plants

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SUMMARY

The *Arabidopsis* NB-LRR immune receptor RPM1 recognizes the *Pseudomonas syringae* type III effectors AvrB or AvrRpm1 to mount an immune response. Although neither effector is itself a kinase, AvrRpm1 and AvrB are known to target *Arabidopsis* RIN4, a negative regulator of basal plant defense, for phosphorylation. We show that RIN4 phosphorylation activates RPM1. RIN4₁₄₂₋₁₇₆ is necessary and, with appropriate localization sequences, sufficient to support effector-triggered RPM1 activation, with the threonine residue at position 166 being critical. Phosphomimic substitutions at T166 cause effector-independent RPM1 activation. RIN4 T166 is phosphorylated in vivo in the presence of AvrB or AvrRpm1. RIN4 mutants that lose interaction with AvrB cannot be coimmunoprecipitated with RPM1. This defines a common interaction platform required for RPM1 activation by phosphorylated RIN4 in response to pathogenic effectors. Conservation of an analogous threonine across all RIN4-like proteins suggests a key function for this residue beyond the regulation of RPM1.

INTRODUCTION

Plants use an active immune system to fend off most microbes. The induction of a successful response to infection relies on specific recognition of pathogen-encoded molecules. Effector proteins are produced by pathogens and translocated into plant cells, where they function as virulence factors. Effectors can be recognized by plant intracellular immune receptors. Type III effectors (T3Es) are produced by Gram-negative phytopathogenic bacteria and injected into host cells through the

hypodermic needle-like type III secretion apparatus (Dodds and Rathjen, 2010; Jones and Dangi, 2006).

Although they can trigger immune receptor function, pathogen-encoded effector proteins, including bacterial T3Es, have evolved to promote virulence (Jakobek et al., 1993). Once delivered, effector proteins are trafficked to various subcellular locations (Nomura et al., 2005). Host-derived modifications such as acylation often influence subcellular targeting of effectors (Nimchuk et al., 2000). Despite their varied sites of action, many effectors share the ability to suppress host defenses via targeting and modification of host proteins that can function to regulate host defense output processes (Gimenez-Ibanez et al., 2009; Hauck et al., 2003; Rosebrock et al., 2007; Shan et al., 2008; Wilton et al., 2010). One example is *Arabidopsis* RIN4, which is a negative regulator of basal plant defense and is targeted by multiple T3Es, including two investigated in this study, AvrRpm1 and AvrB from *Pseudomonas syringae* (Kim et al., 2005b; Mackey et al., 2002).

Plants encode disease resistance proteins that recognize the presence of effectors (Dangi and Jones, 2001; Dodds and Rathjen, 2010; Jones and Dangi, 2006). The majority of intracellular plant disease resistance proteins share a common a central nucleotide binding domain and C-terminal leucine-rich repeats (NB-LRR). The N terminus of RPM1 is composed of a coiled-coil domain (CC-NB-LRR), while a second class of NB-LRR proteins has Toll/IL-1 motifs at their N termini (TIR-NB-LRR). These proteins are analogous to animal innate immune receptors of the NLR class (Ting et al., 2008).

Arabidopsis encodes ~150 NB-LRR proteins, a number that might seem insufficient to offer direct recognition of the diversity of pathogen-encoded effector proteins. However, if pathogen effectors repeatedly target a finite number of host molecules, then NB-LRR proteins indirectly recognizing perturbation of these molecules could provide a robust protective function (Dangi and Jones, 2001; Jones and Dangi, 2006). RIN4 and associated proteins provide evidence for this hypothesis. RIN4 is a negative regulator of immune responses elicited by microbe-associated molecular patterns (MAMPs) (Kim et al., 2005b). Independently evolved T3Es that suppress MAMP-triggered

immunity (MTI) target RIN4; these include AvrRpm1, AvrB, AvrRpt2, HopF2 (Axtell and Staskawicz, 2003; Mackey et al., 2002, 2003; Wilton et al., 2010), and potentially others (Luo et al., 2009). *Arabidopsis* deploys two CC-NB-LRR proteins, RPM1 and RPS2, to monitor RIN4 integrity (Axtell and Staskawicz, 2003; Mackey et al., 2002, 2003). Soybean and lettuce deploy additional NB-LRR-proteins that likely monitor RIN4 orthologs (Ashfield et al., 2004; Jeuken et al., 2009). RPM1 and RPS2 interact with RIN4 at the plasma membrane in unchallenged *Arabidopsis* (Axtell and Staskawicz, 2003; Mackey et al., 2002, 2003). AvrRpm1, AvrB, and AvrRpt2 are acylated subsequent to delivery and thus localized to the host plasma membrane, where they encounter their targets, including RIN4 (Nimchuk et al., 2000). RPM1 responds to AvrRpm1 and AvrB, both of which interact with and induce phosphorylation of RIN4. RPS2 responds to AvrRpt2, a cysteine protease effector that cleaves RIN4 at two sites (Chisholm et al., 2005; Coaker et al., 2005; Kim et al., 2005a). Strong “effector-triggered immunity” (ETI) is induced upon NB-LRR activation. This response is sufficient to bypass blocks in the MTI output caused by other codelivered effectors and leads to an efficient plant immune response.

Our current model holds that RPM1 indirectly recognizes AvrRpm1 and AvrB via RIN4 phosphorylation (Mackey et al., 2002). Thus, we sought to identify phosphorylation sites and other RIN4 residues that regulate RPM1 function. RIN4 has two plant-specific NOI domains (Pfam: PF05627). The C-terminal NOI (NOI2) includes amino acids 142–176, which were cocrystallized with AvrB and contain the AvrB binding site (BBS) (Desveaux et al., 2007). We show here that NOI2 is necessary and, together with the C-terminal RIN4 acylation site required for proper membrane targeting (Kim et al., 2005a), sufficient for effector-triggered RPM1 function. Point mutation of RIN4 residues in this domain revealed that T166 is necessary for AvrB-triggered RPM1 activation. Phosphomimic substitutions at T166 caused effector-independent RPM1 activation, which, like effector-triggered RPM1 activation, is dependent on the RPM1 P loop. RIN4 T166 is phosphorylated in vivo in the presence of AvrB or AvrRpm1. A RIN4 T166A mutant that cannot be phosphorylated fully disrupts AvrB activation of RPM1 and partially disrupts AvrRpm1 activation of RPM1, indicating that AvrRpm1 and AvrB have overlapping but distinguishable mechanisms of activating RPM1. Additional mutations in residues around T166 compromise the ability of RIN4 to interact with both AvrB and RPM1, indicative of a common interaction platform. We conclude that effector-dependent phosphorylation of RIN4 T166 activates RPM1.

RESULTS

The C-Terminal NOI2 Domain of RIN4 Is Sufficient to Trigger RPM1-Mediated Hypersensitive Response

RIN4₁₄₂₋₁₇₆ mediates AvrB interaction (Desveaux et al., 2007). This short region, which includes one of the two AvrRpt2 cleavage sites in RIN4 (RCS2) (between positions 152 and 153), is part of the NOI2 domain conserved in RIN4 homologs from mosses to all flowering plants analyzed to date. We constructed two RIN4 deletion mutants (Figure 1A). The first (1Δ141) expresses NOI2 and the C-terminal palmitoylation/prenylation sequence required for RIN4 localization (Kim et al.,

2005a). The second (149Δ176) deletes the BBS/NOI2 domain. The former construct tests for sufficiency of this domain in RIN4 function, while the latter tests for its necessity. We expressed these derivatives and a wild-type RIN4 control from the native *RIN4* promoter and coding sequence with N-terminal T7 epitope tags as transgenes in *RPM1::myc rpm1 rps2 rin4* (shortened to *RPM1::myc r1 r2 r4* in some figures) (see Experimental Procedures). Homozygous T3 lines expressed RIN4 protein of the appropriate apparent molecular weight (Figure 1B). As expected, *RPM1::myc rpm1 rps2 rin4* is effectively *rpm1*, since RIN4 is required for RPM1 accumulation and hence function (Figures 1C and 1D) (Mackey et al., 2002). RIN4 1Δ141 complemented AvrB- and AvrRpm1-triggered RPM1 function as well as the full-length *RIN4* cDNA transgene (FL:RIN4). By contrast, RIN4 149Δ176 did not. These results were confirmed using conductivity measurements of ion leakage (hereafter “conductivity”) as a proxy for cell death (Figure 1D). In pathogen growth restriction assays (Figure 1E), RPM1 function was restored in transgenic lines expressing FL:RIN4 or RIN4 1Δ141, but not in those expressing RIN4 149Δ176. These data were consistent with RPM1-myc accumulation levels observed in the respective lines (Figure 1F). Thus, RIN4₁₄₉₋₁₇₆ is necessary and, in the presence of required localization sequences, sufficient to mediate RPM1 accumulation and effector-triggered function.

RIN4 Residues Contacting AvrB Are Required for Interaction

We generated missense mutants in the BBS based on contact residues in the cocrystal structure between RIN4₁₄₉₋₁₇₆ and AvrB (Figure S1A) (Desveaux et al., 2007). Yeast two-hybrid data confirmed that RIN4 142Δ176 failed to interact with AvrB (Figure S1B). Further, mutation of I168A and F169A in the RIN4 BBS disrupted interaction with AvrB (Figure S1C). Interestingly, a RIN4 T166A mutant retained—and RIN4 T166D, a phosphomimic mutant, lost—interaction with AvrB. Expression of all RIN4 mutants was confirmed by immunoblot in total yeast protein extracts after mating (Figure S1D); hence, loss of interaction with AvrB is due to RIN4 mutation.

RPM1- and RIN4-Dependent, AvrB-Triggered Hypersensitive Response Reconstituted in *Nicotiana benthamiana*

We optimized a heterologous *Agrobacterium*-mediated transient assay system in *N. benthamiana* to test whether the RIN4 BBS mutants affect the AvrB-elicited, RPM1-mediated hypersensitive response (HR). HR in *N. benthamiana* was observed visually and by trypan blue staining and was quantified by conductivity assays. At the optimized concentrations of each strain, no single or two-partner coinfiltrations of AvrB, RIN4, or RPM1 resulted in cell death (Figure S2A). An AvrB G2A mutant, which is mislocalized due to the lack of a required myristoylation site, did not induce HR, consistent with previous data from *Arabidopsis* (Nimchuk et al., 2000). Leaves infiltrated with AvrB, RIN4, and RPM1 showed onset of conductivity 5–8 hr and macroscopic HR 12 hr after estradiol treatment; no observable phenotype was detected in the other infiltrations (Figures S2A and S2B). Overexpression of RPM1 did not result in ectopic cell death (Figure S2D). Overexpressed AvrB caused cell death at OD₆₀₀ = 0.05 and above, indicating that it is active in *N. benthamiana* (Kang et al., 2010; Schechter et al., 2004). We

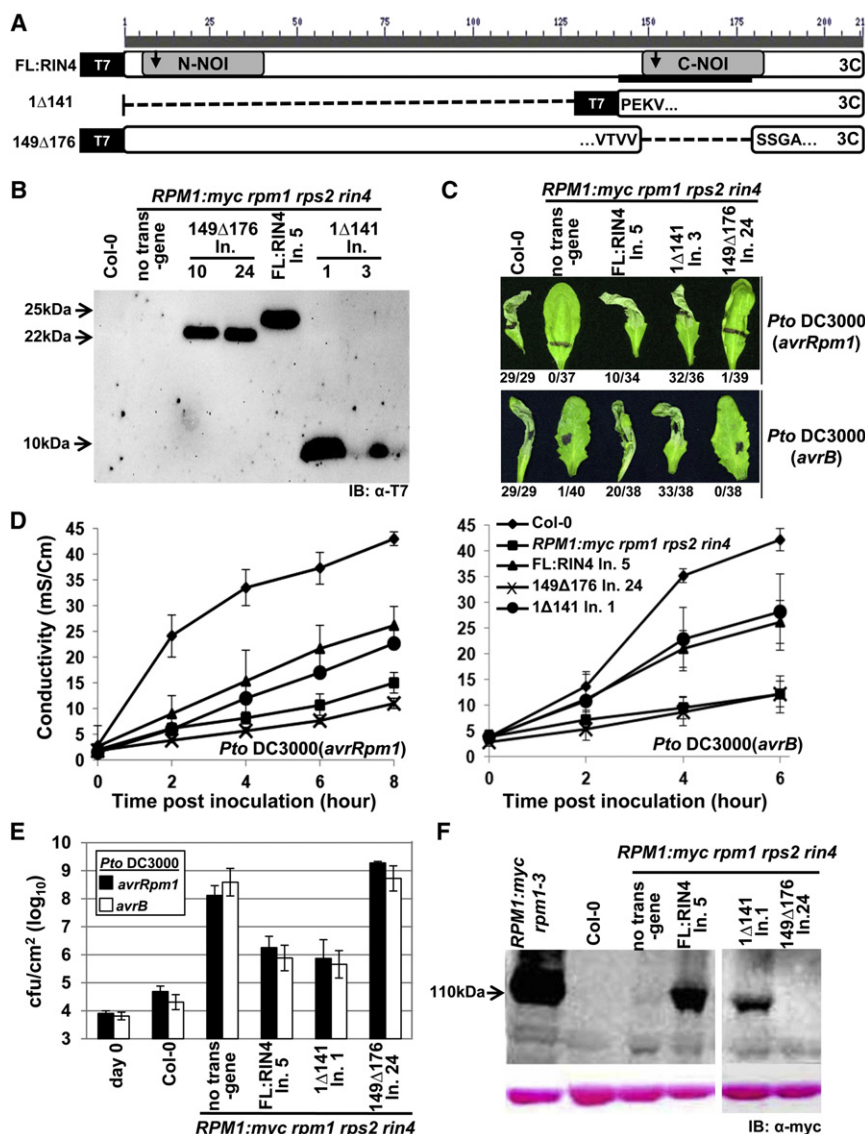


Figure 1. The C-Terminal NOI of RIN4 Is Required for RPM1 Function

(A) Schematic diagram of RIN4 derivatives. Gray boxes are N- and C-terminal NOI domains, the black bar is the AvrB binding site (BBS), the arrows indicate the two AvrB cleavage sites, and the "3C" represents the C-terminal palmitoylation/prenylation site (Kim et al., 2005a). Within the derivatives, the amino acids flanking the break-points are indicated. Each derivative has an N-terminal T7 epitope tag.

(B) α-T7 immunoblot of microsomal membrane protein fractions from transgenic lines expressing the indicated RIN4 derivatives from (A) under control of the native *RIN4* promoter in *RPM1:myc rpm1 rps2 rin4* plants. The background pattern differs in the right panel because this is a higher percentage gel used to resolve 1Δ141 (9 kDa). Line numbers designate plant families homozygous for a single insertion locus that were derived from independent T-DNA insertion events.

(C) HR phenotypes of the indicated plants after infiltration with 5×10^7 cfu/ml of *Pto* DC3000 expressing AvrRpm1 or AvrB, as noted at right. Representative leaves were photographed 20 hr after infiltration, and the numbers below indicate the occurrence of macroscopic HR per number of tested leaves.

(D) Conductivity measurements after infiltration of the indicated plants with 5×10^7 cfu/ml of *Pto* DC3000 expressing AvrRpm1 (left) or AvrB (right). Eight-leaf discs that received the same infiltration were floated in a single tube, and the conductivity of the solution was measured over time. Standard errors are from data combined from three separate experiments.

(E) Growth analysis 3 days after infiltration of 10^5 cfu/ml of *Pto* DC3000 expressing AvrRpm1 or AvrB into the indicated plants. The day 0 measurements show the number of bacteria in Col-0 plants immediately following infiltration. The results are from one of four representative experiments. Standard errors are from three separate experiments.

(F) RPM1 expression in microsomal fractions from the indicated lines. The strong signal in the line *RPM1:myc rpm1-3* shows the high level of RPM1:myc accumulation in the presence of native RIN4.

avoided this background by lowering the amount of infiltrated *Agrobacterium* cells. At $OD_{600} = 0.02$; no visible phenotype was observed, though there was detectable expression of AvrB protein (Figure S2E). Thus, we infiltrated *Agrobacteria* at $OD_{600} = 0.02$ for AvrB, 0.4 for RIN4, and 0.4 for RPM1 for all further experiments, including the final data displayed in Figure S2A demonstrating specific reconstitution of RPM1- and RIN4-dependent, AvrB-triggered HR.

We analyzed the function of our RIN4 BBS mutants in this *N. benthamiana* system. Constructs that expressed RIN4 H167A supported AvrB-triggered, RPM1-dependent HR, but RIN4 derivatives I168A and F169A, an HIF-AAA triple mutant, and a mislocalized nonfunctional AvrB G2A did not (Figures 2A and 2C). These results mirrored those from yeast two-hybrid experiments (Figure S1). RIN4 T166A, which retained interaction with AvrB

(Figure S1), lost the ability to support AvrB-triggered, RPM1-dependent HR (Figure 2B). On the other hand, RIN4 T166D, which cannot interact with AvrB (Figure S1), supported RPM1-dependent HR, even in the absence of AvrB or in the presence of AvrB G2A (Figure 2B; summarized in Table S1). Thus, a RIN4 T166D phosphomimic mutant renders RPM1 activation AvrB independent, indicating that this residue might be phosphorylated as part of the normal AvrB-triggered activation of RPM1. Equal protein expression was confirmed by immunoblotting (Figure 2D).

AvrB-Independent Activation of RPM1 on Membranes Can Be Driven by RIN4 Phosphomimics and Requires a Conserved RPM1 P Loop Residue

Only RIN4 T166D activated RPM1 in the absence of AvrB (Figures 3A and 3B). Neither RIN4 T166D nor any other RIN4

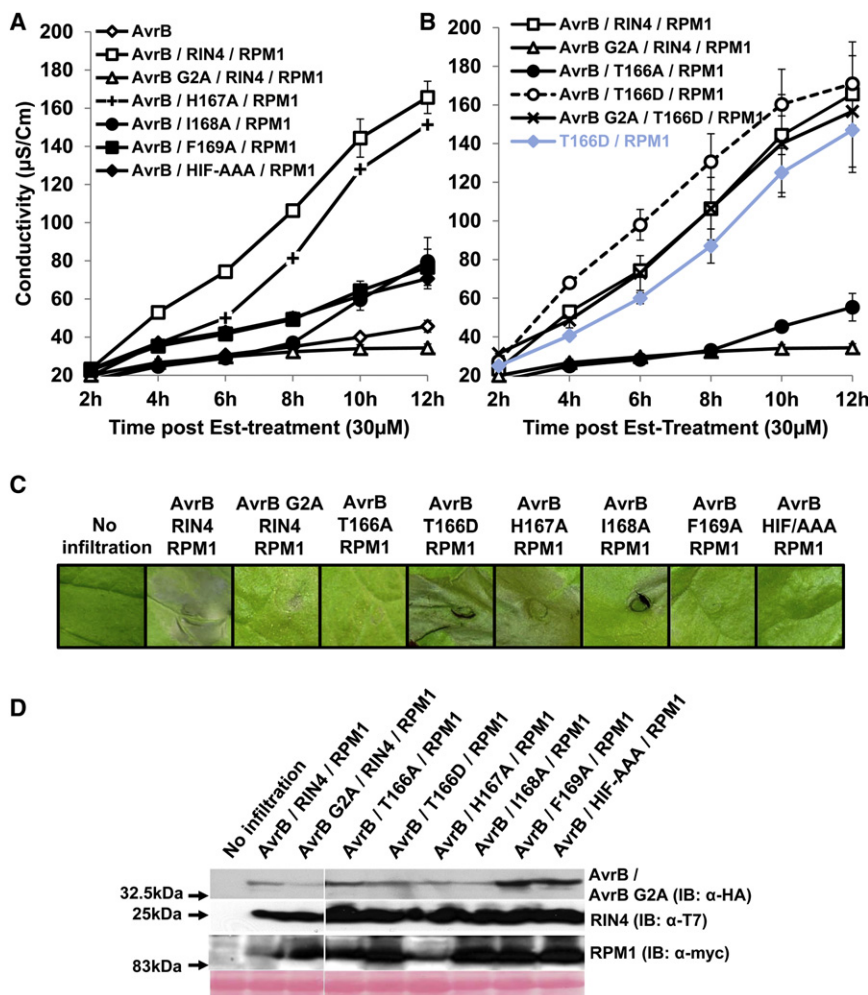


Figure 2. RIN4 T166 Is Required for AvrB-Mediated, RPM1-Dependent HR in *Nicotiana benthamiana*, and a Phosphomimic of this Residue Confers Effector-Independent RPM1 Activation

(A) Conductivity measurements after agro-infiltration with strains expressing the indicated proteins. *N. benthamiana* leaves were hand-infiltrated with *Agrobacterium* C58C1 strains expressing AvrB/AvrB G2A, RIN4, H167A, I168A, F169A, HIF-AAA mutant, and RPM1 as described in Figure S2A. 30 μM of Est was applied 2 days after coinfiltration. Some error bars are smaller than the symbols.

(B) Coinfiltration of AvrB and RPM1 with RIN4 T166A and T166D mutants. This result was obtained from the same experiments in (A). Error bars in (A) and (B) represent 2 × SEM. These results were confirmed four times.

(C) Visible phenotypes of infiltrated *N. benthamiana* leaves. Two independent leaves were infiltrated with the indicated constructs. One leaf was used to take the picture for phenotypes and the second leaf was used to extract proteins for immunoblot in (D). Pictures were taken 12 hr after Est treatment. The result is one of four replicates. (D) Immunoblots with α-HA, α-T7, and α-myc for AvrB/AvrB G2A, RIN4/BBS mutants, and RPM1, respectively, following *Agrobacterium*-mediated transient expression. Protein samples were harvested 6 hr after Est treatment.

BBS mutant caused HR in the absence of RPM1 (Figure S3). We extended our finding that RIN4 T166D drives effector-independent RPM1 activation using RIN4 T166E, with glutamic acid an alternative phosphomimetic residue (Figure 3B). We demonstrated that RIN4 T166K does not cause HR, demonstrating specificity for phosphomimetic alterations rather than mere charge change (Figure 3B). RPM1 and all RIN4 T166 derivatives were expressed in the *Agrobacterium*-mediated transient assay (Figure 3C). These data, together with the loss of HR observed following coexpression of AvrB/RIN4 T166A/RPM1 (Figure 2), strongly suggest that RIN4 T166 is phosphorylated in response to AvrB and that this modification is necessary for subsequent RPM1 activation.

RPM1 and RIN4 are both associated with the plasma membrane (Boyes et al., 1998; Kim et al., 2005a). Both RIN4 T166D and RPM1 were detected in microsomes from transiently expressing *N. benthamiana* leaves (Figure 3D). RPM1, RIN4, and RIN4 T166D were all enriched in plasma membrane fractions following two-phase partitioning (Figure 3E) (Boyes et al., 1998), indicating that RPM1 activation via RIN4 T166D occurs there. Further, both RIN4 and RIN4 T166D can be coimmunoprecipitated with RPM1 in vivo from microsomes (see below). Thus, RIN4 T166D associates with and likely modulates the activity of RPM1 on the plasma membrane.

NB-LRR activation (Takken and Tameling, 2009; van Ooijen et al., 2007). The RPM1 G205E mutation in the P loop exhibited a loss-of-function phenotype (Tornero et al., 2002). We used this allele to address whether RIN4 T66D-driven, RPM1-dependent HR requires a wild-type P loop. RIN4 T166D-mediated activation of RPM1 G205E is significantly impaired (Figure 3F). Thus, activation of RPM1 by RIN4 T166D is regulated by canonical P loop function, similar to the requirements for activation of RPM1 by AvrB and AvrRpm1 during infection of *Arabidopsis* (Tornero et al., 2002).

The RIN4 T166D Phosphomimic Retains the Ability to Be Cleaved by AvrRpt2 in *N. benthamiana*

RIN4 is a target for a third *P. syringae* T3E protein, the cysteine protease AvrRpt2. Cleavage of RIN4 at the second of two specific sites activates RPS2-mediated defense resistance (Mackey et al., 2003; Kim et al., 2005a). Coexpression of RIN4 and AvrRpt2 in *Agrobacterium*-mediated *N. benthamiana* transient assays results in RIN4 cleavage (Day et al., 2005). Both RIN4 and RIN4 T166D were cleaved by AvrRpt2 but not by an AvrRpt2 catalytic mutant (C122A) in this assay (Figure S4). Thus, a phosphomimic of RIN4 on T166 cannot block cleavage by AvrRpt2.

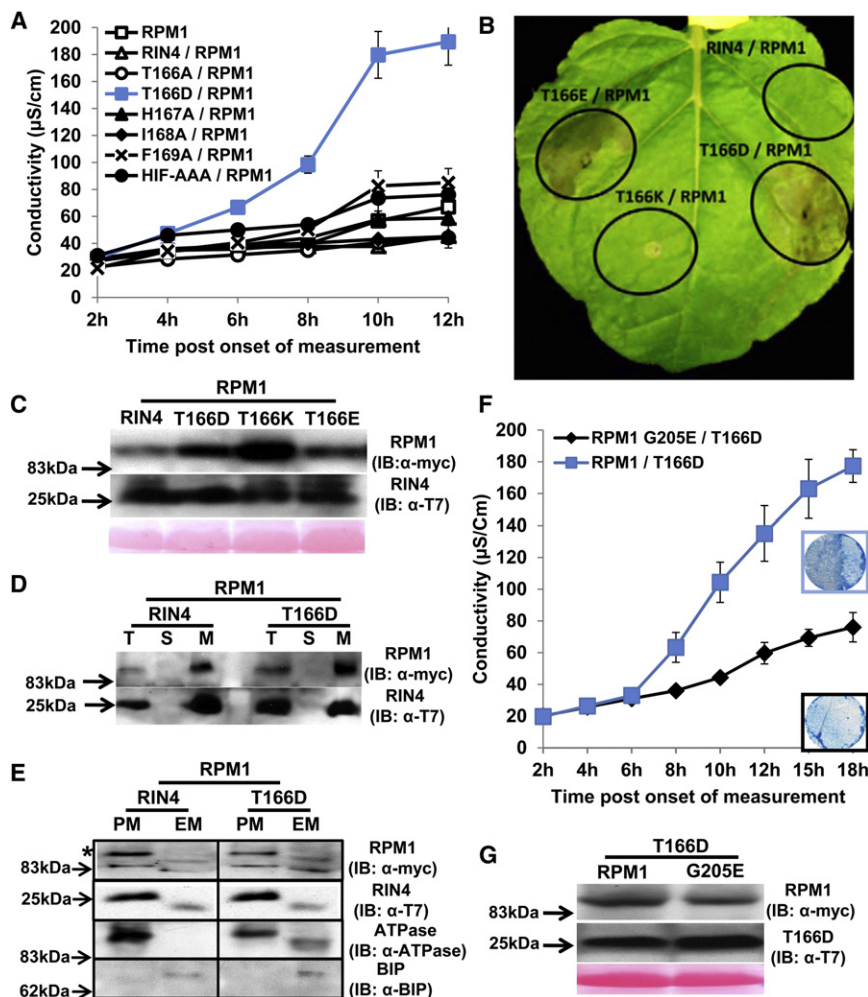


Figure 3. RIN4 T166D Activity Is Dependent on RPM1 P Loop Function in *Nicotiana benthamiana*

(A) The phosphomimic RIN4 T166D mutant drives effector-independent, RPM1-mediated HR. Conductivity measurements were performed with *N. benthamiana* leaves infiltrated with *Agrobacterium* C58C1 strains expressing RIN4 BBS mutants ($\text{OD}_{600} = 0.4$) and RPM1:myc ($\text{OD}_{600} = 0.4$). The measurements began 2 days postinfiltration. Experiment was repeated three times with similar results. The error bars represent $2 \times \text{SEM}$.

(B) Phenotypes of RIN4 T166 derivatives. RIN4 T166D, RIN4 T166E, and RIN4 T166K driven by the RIN4 native promoter were coinfiltrated as in (A). Photo was taken 3 days after coinfiltration.

(C) Expression of RPM1 and RIN4 T166 derivatives. RPM1 and RIN4 T166 derivatives were expressed, and variation does not account for the observed phenotypes. Immunoblots with α -myc and α -T7 were performed with 2-day-old samples postinfiltration.

(D) The RIN4 phosphomimic T166D is localized to a microsomal fraction. *N. benthamiana* leaves were coinfiltrated as in (A). Proteins were extracted from leaf tissues at the onset of HR from T166D/RPM1 coinfiltration, which corresponded to 8 hr in the conductivity experiment in (A). Results were repeated twice. Total (T), soluble (S), and microsomal (M) fractions were loaded at a 1:1:5 ratio, followed by immunoblotting with α -T7 and α -myc to detect RIN4 and RPM1, respectively.

(E) Two-phase partitioning of RIN4 and RPM1. RIN4 and RIN4 T166D mutant were coinfiltrated with RPM1 as described in (D). The microsomal extraction was used as input for two-phase partitioning. The upper fraction, for plasma membrane (PM), and the lower fraction, for endomembranes (EM), were loaded at equal yield, followed by immunoblotting with α -myc and α -T7 to detect RPM1(*) and RIN4, respectively. PM-localized ATPase and EM-localized BIP proteins represented the efficiency of fractioning for PM and EM.

(F) Conductivity measurements and HR phenotype after coinfiltration of RIN4 T166D with either RPM1 or an RPM1 G205E. *N. benthamiana* leaves were hand-infiltrated with *Agrobacterium* C58C1 strains expressing T166D mutant ($\text{OD}_{600} = 0.4$) and either *pRPM1:RPM1:myc* ($\text{OD}_{600} = 0.4$) or RPM1:myc G205E ($\text{OD}_{600} = 0.8$). C58C1 was used as filler to make up the difference in OD between RPM1:myc and RPM1:myc G205E with $\text{OD}_{600} = 0.4$. The measurements started 2 days postinfiltration. This result was one of two repeats. Trypan blue staining with leaf discs covering half of the infiltrated zone was performed 2.5 days after infiltration indicated 12 hr in conductivity measurement.

(G) Expression of RPM1:myc and RPM1:myc G205E. Protein samples from (F) were prepared 2 days postinfiltration. The immunoblot was performed with α -myc.

RIN4 T166 Contributes to but Is Not Essential for AvrRpm1-Dependent, RPM1-Mediated HR in *N. benthamiana*

AvrRpm1 is a *P. syringae* T3E unrelated to AvrB that can also activate RPM1-mediated HR and be coimmunoprecipitated with RIN4 (Mackey et al., 2002). AvrRpm1 does not interact with RIN4 in Y2H, and it is unstable and unstructured in vitro following purification (K.C. and J.L.D., unpublished data). Therefore, the nature of its direct interaction with RIN4, if any, remains elusive. Hence, we sought to functionally cross-reference the RIN4 residues required for AvrB-triggered RPM1 activation to AvrRpm1.

We reconstituted a functional AvrRpm1-triggered RPM1 activation assay in *N. benthamiana* (see Experimental Procedures). We observed that RIN4 I168A and F169A mutants could not support AvrRpm1-triggered, RPM1-dependent HR (Figure 4), consistent

with their phenotypes in AvrB-triggered, RPM1-dependent HR (Figure 2). Wild-type RIN4 and, to a slightly lesser extent, RIN4 T166A supported RPM1-dependent, effector-induced conductivity (Figure 4A) and HR (Figure 4B). Protein expression for AvrRpm1, RIN4, and RPM1 was confirmed (Figure 4C). These data, combined with those in Figure 2, indicate that RIN4 T166 is required for AvrB-triggered, RPM1-dependent HR, but not essential for AvrRpm1-triggered, RPM1-dependent HR in *N. benthamiana*.

Native Expression Level RIN4 T166D Transgenic Lines Exhibit Ectopic Basal Defense Phenotypes

We recapitulated the key results from our transient expression system in transgenic *Arabidopsis* plants. All native promoter RIN4 constructs used for *Agrobacterium*-mediated transient assay on *N. benthamiana* were stably transformed into RPM1:myc

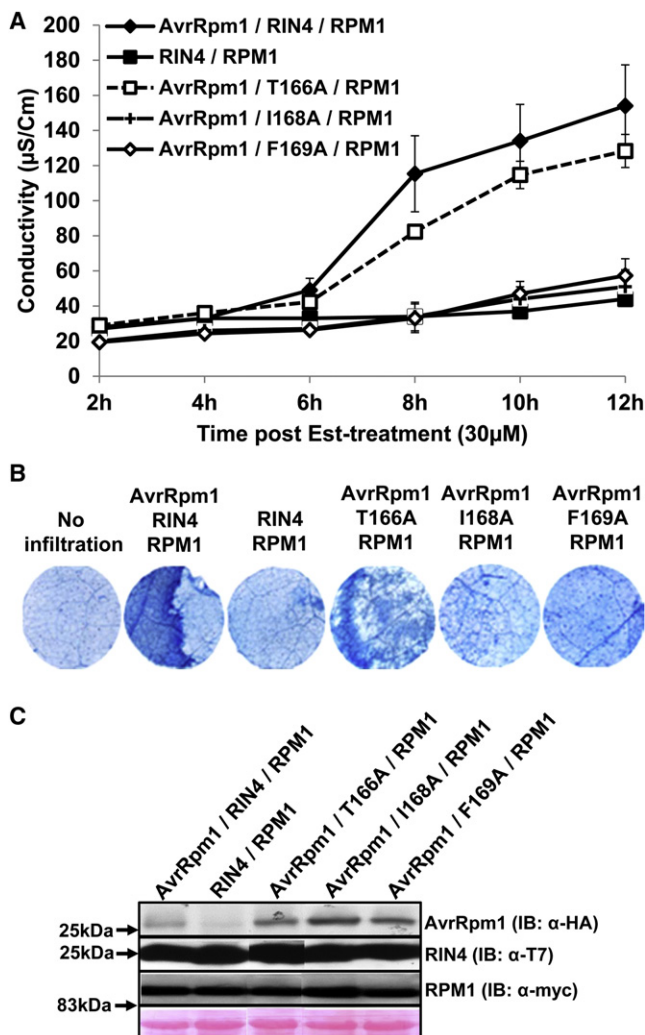


Figure 4. RIN4 T166 Contributes to AvrRpm1-Dependent, RPM1-Mediated HR in *N. benthamiana*

(A) Conductivity measurements after agro-infiltration with strains expressing the indicated proteins. *N. benthamiana* leaves were hand-infiltrated with Agrobacterium C58C1 strains as in Figure 2A except *Est:AvrRpm1-HA* ($OD_{600} = 0.1$) instead of *Est:AvrB:HA*. Coinfiltration of RIN4 and RPM1:myc was used as a negative control with C58C1 cells ($OD_{600} = 0.1$). The result was repeated three times. Measurement started 2 hr postinduction with 30 μM Estradiol. Error bars represent 2× SEM.

(B) HR phenotypes of infiltrated *N. benthamiana* leaves. Trypan blue staining was performed with leaf discs that covered half of an infiltrated zone at 8 hr after Est treatment. Data represent one of three independent experiments with consistent results.

(C) Immunoblots with α-HA, α-T7, and α-myc to detect AvrRpm1, RIN4, and RPM1, respectively. Protein samples were extracted from leaf tissues harvested 6 hr after Est treatment.

rpm1 rps2 rin4. We obtained at least two independent homozygous T3 transgenic lines expressing each RIN4 BBS mutant. We observed dwarfism and chlorosis in both independent T166D lines, especially under long day conditions, and no obvious phenotype in lines expressing the other BBS mutants (Figure S5A). Each RIN4 BBS mutant protein was expressed at levels approximating that of wild-type RIN4 in Col-0 and *rpm1-3* (Figure S5B).

We noted a mild ectopic cell death in lines expressing RIN4 T166D (Figure S5C). Furthermore, we observed ectopic PR1 protein expression in RIN4 T166D mutants, consistent with the lesion and morphology phenotypes of constitutive defense mutants (Figure S5D). The mild constitutive defense activation phenotype expressed by RIN4 T166D transgenics was sufficient to limit growth of the virulent bacterial pathogen *Pto* DC3000 (Figure S5E). These phenotypes were RPM1 dependent (Figure S5F).

RIN4 T166 Is Essential for AvrB-Triggered RPM1 Function in *Arabidopsis*

We tested RPM1 function following infiltration of *Pto* DC3000 expressing either *avrB* or *avrRpm1* into leaves of the various RIN4 BBS expressing transgenic lines and appropriate controls (Figure 5). RIN4 derivatives I168A, F169A, and HIF-AAA did not support AvrB- or AvrRpm1-triggered HR, while the RIN4 H167A did (Figures 5A and 5B). Importantly, RIN4 T166A did not support either HR or increased conductivity following inoculation with *Pto* DC3000(*avrB*) (Figures 5A and 5B). RIN4 T166A supported an intermediate level of RPM1-dependent HR triggered by *Pto* DC3000(*avrRpm1*) compared to Col-0 or RIN4 wild-type transgenic plants (gRIN4) and negative control plants (*rpm1-3* and *RPM1:myc rpm1 rps2 rin4*) (Figure 5A). We confirmed and quantified this intermediate phenotype in leaves from two independent, homozygous, transgenic lines following inoculation with *Pto* DC3000(*avrRpm1*) (Figure 5B, T166A lines 32.6 and 33.5). These results are consistent with those from the *N. benthamiana* reconstruction system. Finally, we tested RPM1-mediated bacterial growth restriction in the RIN4 BBS mutant lines following low-dose inoculation with *Pto* DC3000(*avrB*) or (*avrRpm1*). Concordant with HR assay results, RIN4 T166A exhibited a loss of RPM1 function phenotype in response to *Pto* DC3000(*avrB*) and slightly reduced function, relative to gRIN4, in response to *Pto* DC3000(*avrRpm1*) (asterisks in Figure 5C). These results indicate that RIN4 T166 is required for AvrB-triggered RPM1 function and contributes to, but is not essential for, AvrRpm1-triggered RPM1 function (summarized in Table S1).

Oddly, the RIN4 T166D transgenic lines exhibited RPM1-dependent HR triggered by AvrB (weak) and AvrRpm1 (intermediate) (Figure 5A) and, in fact, by *Pto* DC3000 (weak) (Figure S6A). These results, coupled with PR1 expression data in Figure S6B, indicate that ectopic RPM1 signaling in RIN4 T166D-expressing lines results in a lowered threshold for activation of the low level of RPM1 that accumulates in these lines (see below).

RIN4 T166 Is Phosphorylated in Response to AvrB and AvrRpm1

We addressed whether T7 epitope-tagged RIN4 T166 could be phosphorylated by immunoprecipitation of RIN4 with α-T7-conjugated agarose beads, followed by immunoblotting with a phosphopeptide-specific antibody raised against a 13 amino acid RIN4 peptide containing phosphothreonine (α-pRIN4) (see Experimental Procedures). To enrich for phosphorylated RIN4 in our transient assay, AvrB or AvrRpm1 and RIN4 or RIN4 T166A mutant were coexpressed without RPM1 (Mackey et al., 2002). As displayed in Figure 6A, signal detected with α-pRIN4 was enriched in α-T7 immunoprecipitates from samples coexpressing wild-type RIN4 with either AvrB or AvrRpm1, compared to samples from extracts coexpressing RIN4 T166A and either effector.

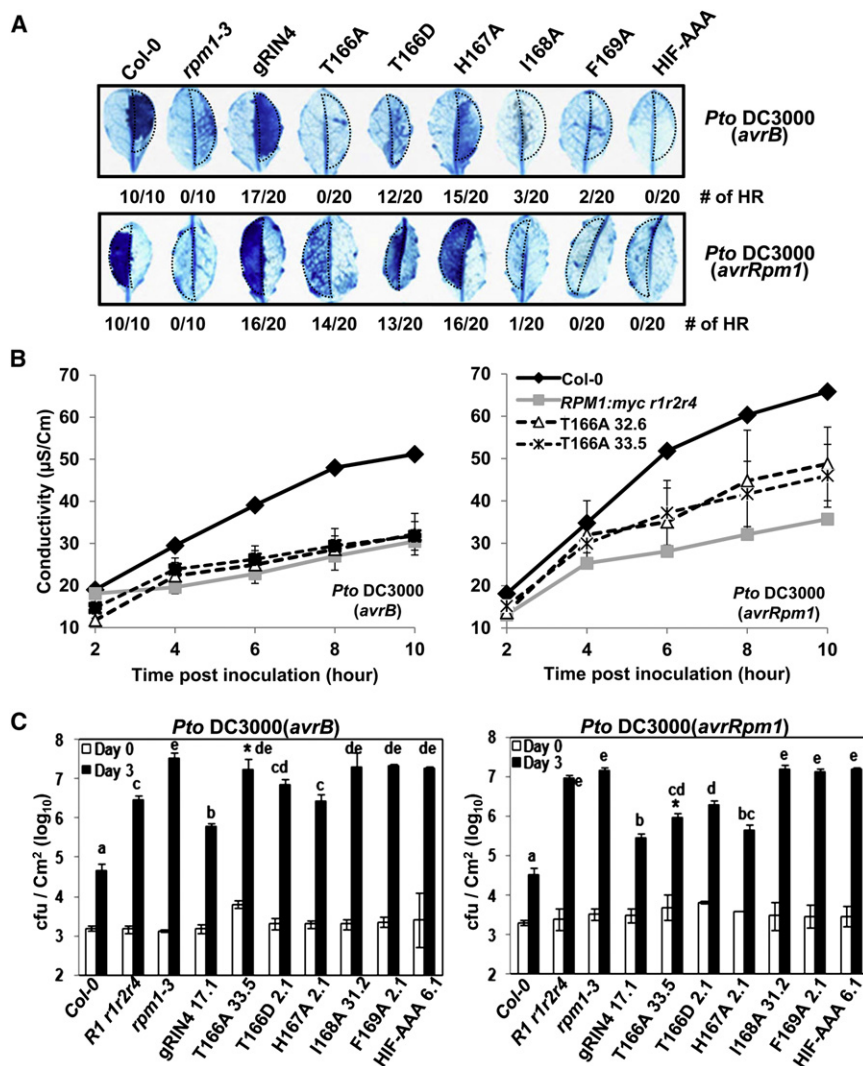


Figure 5. RIN4 T166 Is Required for AvrB-Dependent and Contributes to AvrRpm1-Dependent, RPM1-Mediated HR in Arabidopsis

(A) HR determined by trypan blue staining. Twenty independent leaves from transgenic lines expressing each RIN4 BBS mutant were inoculated with 5×10^7 cfu/ml ($\text{OD}_{600} = 0.1$) of *Pto* DC3000(*avrB*) or (*avrRpm1*) in half of each leaf (dotted area). Leaves were harvested 6 hr after inoculation. The numbers are leaves that displayed the HR phenotype shown over the total. The result was repeated with two independent homozygous transgenic lines for each BBS mutant with similar results.

(B) Conductivity measurements. Two independent homozygous T166A mutant lines and controls shown at right were used to monitor the loss-of-function phenotype with 5×10^7 cfu/ml of *Pto* DC3000(*avrB*) or (*avrRpm1*). Error bar represents $2 \times \text{SEM}$ for RIN4 T166A mutant. Four-leaf discs were used to measure the conductivity of Col-0 and *RPM1:myc rpm1rps2rin4*.

(C) Bacteria growth analysis of *Pto* DC3000(*avrB*) or (*avrRpm1*). Bacteria recovered from infiltrated leaves of each transgenic line indicated or controls at bottom were counted after hand-inoculation with 10^5 cfu/ml for each strain on day 0 and day 3. The result was repeated twice with two independent T3 homozygous transgenic *Arabidopsis* lines from each RIN4 mutant. Error bars represent $2 \times \text{SEM}$. Pairwise comparisons for all means from the day 3 data were performed with one-way ANOVA followed by Tukey-Kramer HSD at 95% confidence limits.

We analyzed RIN4 T166 phosphorylation in transgenic *Arabidopsis*-expressing native levels of either wild-type RIN4 or RIN4 T166A, complementing a *rin4* null allele in the presence *RPM1:myc*. Transgenic plants were infiltrated with *Pto* DC3000 expressing AvrB:HA or AvrRpm1:HA. α -HA and α -T7 immunoblots detected AvrB:HA and AvrRpm1:HA, or the RIN4 derivatives, respectively, in the input for the immunoprecipitations (Figure 6B, top). α -T7 immunoprecipitates were immunoblotted with α -pRIN4 (Figure 6B, bottom). We noted T166-dependent enhancement of α -pRIN4 signal compared to uninfected control in the presence of either effector.

We also demonstrated that the effector-dependent increase in the RIN4 detected with α -pRIN4 is phosphorylation by treating α -T7 immunoprecipitates with calf intestinal phosphatase (CIP), followed by blotting with either α -pRIN4 or α -T7 (Figure 6C). While there is some residual recognition of RIN4 T166A protein by the α -pRIN4 sera, the increased signal it detects is RIN4-pT166. In sum, the results presented in Figure 6 indicate that the presence of either AvrB or AvrRpm1 leads to increased phosphorylation of RIN4 T166 in both *N. benthamiana* and *Arabidopsis* systems.

We therefore performed coimmunoprecipitations with microsomal fractions from the RIN4 BBS mutant transgenic lines (Figure 7). While the RIN4 BBS mutant proteins accumulated equally on microsomes, they supported variable levels of RPM1 accumulation in the input extracts (Figure 7A). Immunoprecipitation of all of the available RPM1 from microsomes led to differentially coimmunoprecipitated RIN4 BBS mutant proteins (Figure 7A). Wild-type RIN4 and RIN4 T166A retained the ability to associate with RPM1 and supported nearly wild-type RPM1 accumulation levels. RIN4 BBS alleles that lost both the ability to interact with AvrB (Figure S1C) and the ability to support AvrB-triggered RPM1 functions (Figure 5) also lost the ability to associate with, and/or support accumulation of, RPM1 (RIN4 I168A, F169A, and HIF-AAA) (Table S1).

The inability of these RIN4 derivatives to support RPM1 accumulation is likely due to a disruption of the interaction between RIN4 and RPM1 at the membrane. This is striking for RIN4 F169A, which fails to coimmunoprecipitate with RPM1. RIN4 T166D drives activation and consequent disappearance of RPM1 at steady state in the transgenics. Nevertheless, a very

RIN4 BBS Residues Are Required for Steady-State Microsomal Accumulation of RPM1

RIN4 can be coimmunoprecipitated with, and is required for accumulation of, RPM1 in unstimulated cells (Belkadir et al., 2004; Mackey et al., 2002). We

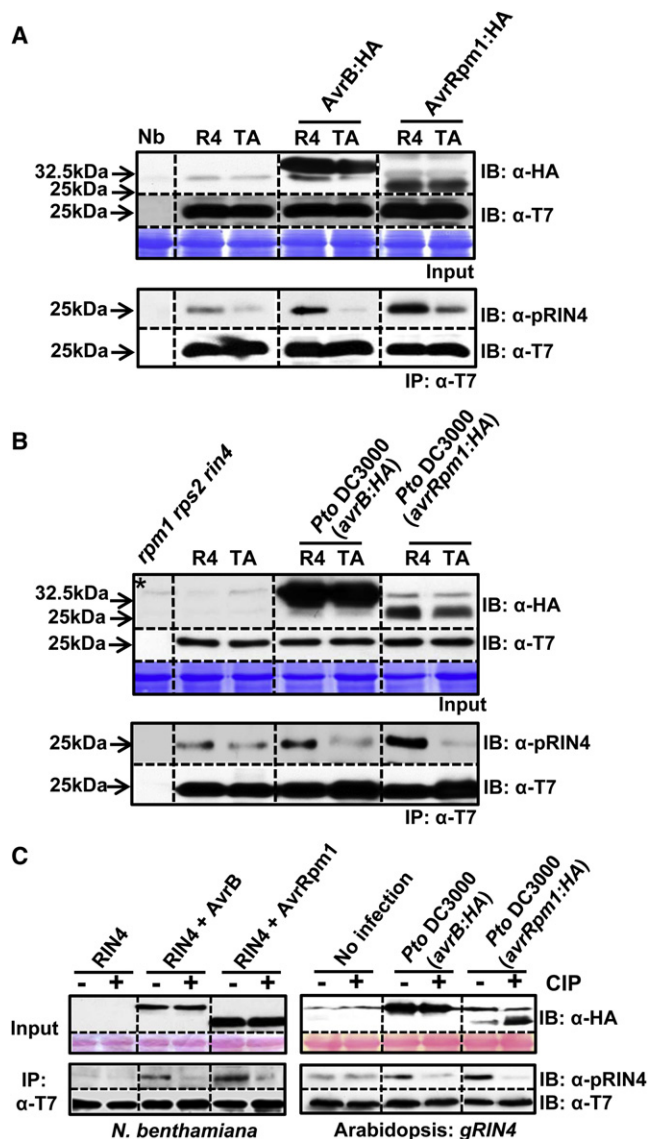


Figure 6. RIN4 T166 Residue Is Phosphorylated by AvrB and AvrRpm1 In Planta

(A) T166-dependent RIN4 phosphorylation in *N. benthamiana*. Immunoprecipitation with α-T7-conjugated agarose beads was used to enrich RIN4 or RIN4 T166A from leaves coinfiltrated with Est:AvrB:HA or AvrRpm1:HA and T7:RIN4 or T7-RIN4 T166A, followed by immunoblotting with α-pRIN4 (phosphopeptide-specific polyclonal antibody) and α-T7. Samples 18 hr after 30 μM Est induction were prepared, and input levels were established by immunoblot with appropriate antibodies (top). α-T7 immunoprecipitates (bottom) were used for immunoblots with α-pRIN4. An immunoblot with α-T7 demonstrated equal expression levels of RIN4 and RIN4 T166A in these immunoprecipitates. The experiment was repeated three times.

(B) RIN4 T166 is phosphorylated in *Arabidopsis* following AvrB or AvrRpm1 delivery from *P. syringae*. Transgenic *Arabidopsis* RIN4 or T166A mutant were inoculated with Pto DC3000(avrB:HA) or (avrRpm1:HA) as described in Figure 5B. Samples were collected 18 hr after infection. Immunoblots and immunoprecipitations were performed as in (A). Asterisk indicates an *Arabidopsis* background band mobility similar to that of AvrB. The data represent one of three experiments with similar results.

(C) The α-pRIN4 antiserum detects phosphorylated RIN4-pT166 in *N. benthamiana* and transgenic *Arabidopsis*. α-T7 immunoprecipitates either from *N. benthamiana* transiently expressing RIN4 and RIN4 with AvrB or AvrRpm1

low level of RPM1 is detected, and it can coimmunoprecipitate RIN4 T166D (Figure 7A). We therefore constructed a RIN4 T166D F169A double mutant. This RIN4 derivative accumulates normally on microsomes, but cannot be coimmunoprecipitated with RPM1 (Figure 7B) or support effector-independent activation of RPM1 HR in the *N. benthamiana* transient assay system (Figure 7C). In this transient expression assay, a high level of RIN4 T166D maintains interaction with the relatively low levels of RPM1, even as the latter is being activated. Hence, RIN4 F169 is required for the interaction of RIN4 with AvrB (Figure S1) and also controls interaction with, and thus stability of, RPM1. Further, this interaction is required for activation of RPM1 by RIN4 T166D.

DISCUSSION

We present a mechanism for effector-dependent activation of a typical NB-LRR plant intracellular immune receptor. *Arabidopsis* RPM1 is activated in response to two unrelated bacterial T3E proteins, AvrB and AvrRpm1. We initially proposed that a host target of both effectors, RIN4, is “guarded” by RPM1. We suggested that modification of RIN4 by AvrB or AvrRpm1 activates RPM1, resulting in suppression of bacterial growth and a HR (Mackey et al., 2002). We noted that the presence of either AvrB or AvrRpm1 resulted in phosphorylation of RIN4, though neither effector has kinase activity in vitro, and we noted that this modification was more pronounced in the absence of RPM1 (Mackey et al., 2002). RIN4 negatively regulates MTI, and both AvrB and AvrRpm1 suppress MTI in plants lacking RPM1 (Kim et al., 2005b). Based on the data presented above, a reasonable speculation is that phosphorylation of RIN4 T166 potentiates the negative regulation of MTI by RIN4. In the absence of RPM1, AvrB or AvrRpm1 “lock” RIN4 as a negative regulator of MTI. RPM1 responds to the effector-induced phosphorylation of RIN4.

The specific RIN4 residues phosphorylated in the presence of AvrB or AvrRpm1 were previously unknown, and a requirement for RIN4 modification in RPM1 activation had not been demonstrated. Here, we provide evidence that phosphorylation of RIN4 T166 is required for AvrB-dependent activation of RPM1 and contributes to AvrRpm1-dependent RPM1 activation. Further, a phosphomimic at this residue (T166D) causes effector-independent activation of RPM1. These data, together with previous publications, provide a mechanism whereby AvrB enters the cell; is targeted by acylation to the host plasma membrane (Nimchuk et al., 2000); is activated (Desveaux et al., 2007), perhaps by a host MAPK (Cui et al., 2010) or other kinases; and enhances the phosphorylation of RIN4 on T166 and potentially other residues. Because AvrB and RPM1 require the same binding site on RIN4, RIN4 phosphorylation is unlikely to occur while it associates with RPM1. AvrB likely dissociates once RIN4 is phosphorylated since the T166D derivative of RIN4 no longer interacts with AvrB. Dissociation of phosphorylated RIN4 from AvrB appears key to RPM1 activation.

(left) or from transgenic *Arabidopsis*, uninfected or infected with Pto DC3000 (avrB:HA) or (avrRpm1:HA) (right), were divided in half to treat calf intestinal phosphatase (CIP). Tissue samples were prepared as in (A) for *N. benthamiana* and (B) for *Arabidopsis*.

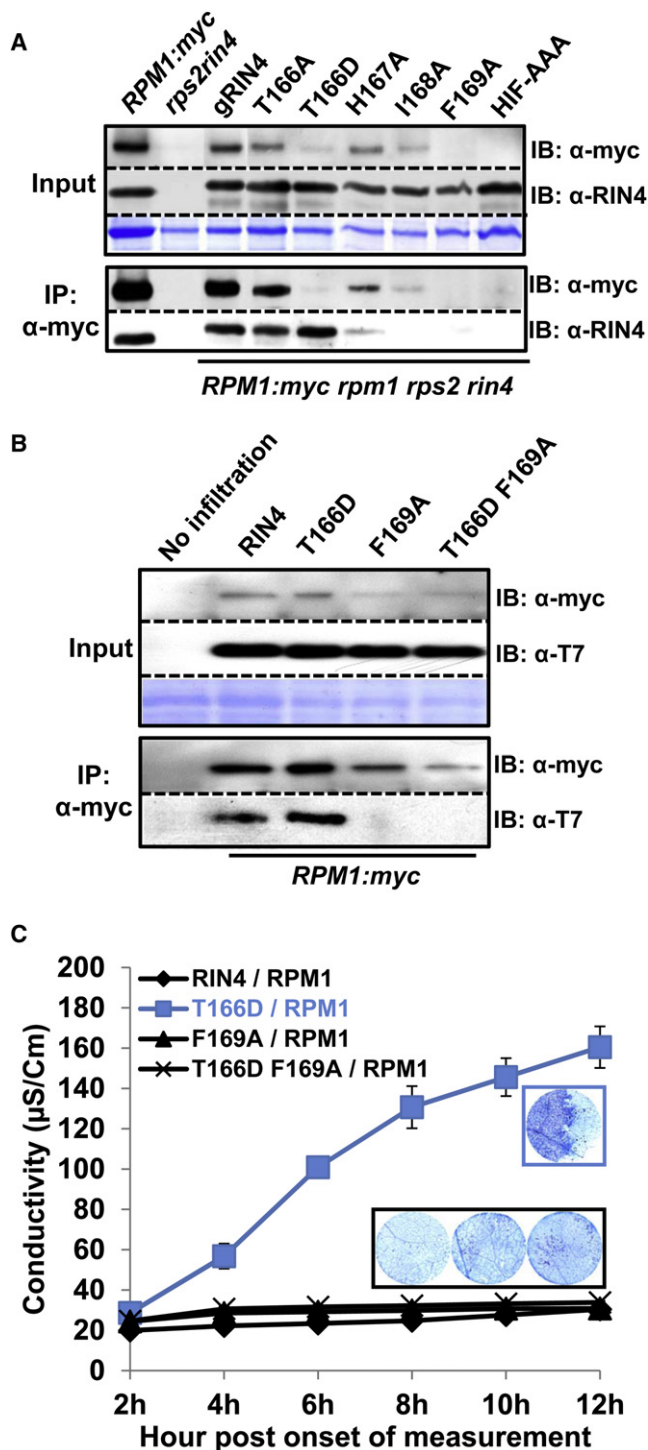


Figure 7. Differential Coimmunoprecipitation of RPM1 with RIN4 BBS Mutants Identifies Residues Required for Interaction and RPM1 Accumulation

(A) Coimmunoprecipitation of RIN4 BBS mutants with RPM1. The microsomal fraction was enriched in extracts from each RIN4 BBS mutant transgenic *Arabidopsis*, followed by immunoprecipitation with α -myc. The overall level of RPM1 is displayed in the input (left top). RIN4 expression in each mutant was confirmed by immunoblotting with α -RIN4. Immunoprecipitated RPM1 was shown by immunoblotting with α -myc. Coimmunoprecipitated RIN4

RIN4 is phosphorylated in the absence of effectors (Mackey et al., 2002) and on residues other than T166 after perception of the flagellin MAMP peptide, flg22 (Nühse et al., 2007). Given the sensitivity of NB-LRR activation, it may be that a threshold level of RIN4 T166 phosphorylation must be attained for RPM1 activation. Additional effector-induced modifications of RIN4, perhaps other phosphorylation events or conformational changes, may increase the propensity of modified RIN4 to activate RPM1. This appears to be the case for AvrRpm1; RIN4 T166A only partially compromises activation of RPM1. Additional phosphorylation of RIN4 by AvrRpm1 is consistent with AvrRpm1 inducing a significantly greater mobility shift in RIN4 than does AvrB (Mackey et al., 2002). Other possible target residues for phosphorylation within the genetically defined region of RIN4 required for AvrRpm1-dependent RPM1 activation include S160 and S161. However, phosphomimics of these residues did not result in effector-independent RPM1 activation, and mutations to alanine did not compromise either AvrRpm1- or AvrB-dependent RPM1 activation. AvrRpm1 may direct functionally relevant phosphorylation or additional modifications of RIN4 residues outside of the NOI2 domain.

Effector-independent activation of RPM1 mediated by RIN4 T166D requires the P loop within the RPM1 NB domain. Hence, pRIN4 T166 is a physiological elicitor of RPM1. Current models of NB-LRR activation envisage an ADP-bound resting state conformation involving intra- and possibly intermolecular interactions that result in the LRR domain inhibiting activation at the NB. Activation is proposed to be driven or accompanied by nucleotide exchange and/or hydrolysis, which are thought to activate downstream processes (Takken and Tameling, 2009; van Ooijen et al., 2007). It has been difficult thus far to establish an order of events for this activation with respect to nucleotide binding and/or turnover. Our results are consistent with a model wherein RPM1 recognition of RIN4pT166 precedes or is coincident with ADP/ATP exchange/hydrolysis, since a loss of function RPM1 P loop mutation also blocks both effector- and RIN4 T166D-mediated RPM1 activation.

Our data support a model in which effector-dependent modification of RIN4 activates RPM1. This model differs from the model of activation of RPS2 via elimination of RIN4 that we and others proposed (Axtell and Staskiewicz, 2003; Mackey et al., 2003). RIN4 is, genetically, a negative regulator of both RPM1 and RPS2 (Belkhadir et al., 2004). However, in the absence of RIN4, ectopic activation of RPS2 occurs, and the result is seedling lethality. In contrast, the lack of RIN4 contributes only weakly to ectopic RPM1 activation (Belkhadir et al., 2004). The inability of RIN4 T166D F169A to activate RPM1 indicates that RIN4 must interact with RPM1 to activate it and that merely disrupting the association of RPM1 with RIN4 is insufficient to fully activate RPM1.

with RPM1 was confirmed with α -RIN4 immunoblot. Two-week-old seedlings from each line were used to collect the microsomal fraction.

(B) Loss of coimmunoprecipitation of RIN4 T166D F169A with RPM1. Agrobacterium transient assays were performed as in Figure 3. Loading controls, immunoprecipitation with α -myc, and subsequent immunoblots were performed as in (A), with the use of α -T7 to detect RIN4 and RIN4 BBS mutants. (C) Loss of effector-independent RPM1 activation in RIN4 T166D F169A. Agrobacterium transient assays, conductivity measurements, and trypan blue staining were performed as in Figure 3.

It is instructive to compare the regulation of RPM1 and RPS5 activation via RIN4 to other well-studied examples of recognition of modified self by plant NB-LRR proteins. The *Arabidopsis* RPS5 NB-LRR protein is activated by cleavage of the host kinase PBS1 by the T3E cysteine protease AvrPphB (Ade et al., 2007; Shao et al., 2003). There is no ectopic RPS5 activation in *pbs1* null plants, indicating that PBS1 is not formally a negative regulator of RPS5. However, AvrPphB suppresses MTI by cleaving PBS1 and related host kinases that may function redundantly to inhibit RPS5 activation (Zhang et al., 2010). Similarly, the Pto kinase family in tomato is targeted by multiple T3Es, and post-translational modification of these kinases activates the Prf NB-LRR protein in ETI (Ntoukakis et al., 2009). Thus, cleaved PBS1 and modified Pto are likely to activate RPS5 and Prf, respectively, similar to the activation of RPM1 by phosphorylated RIN4. The activation of plant NB-LRR proteins by modified self may be similar to the activation of animal NLR proteins of similar structure in response to the presence of MAMPs and/or nonself (Vance et al., 2009).

RIN4 is targeted by four different bacterial T3Es that perturb it in four different ways: proteolysis by AvrRpt2 (Axtell et al., 2003; Axtell and Staskawicz, 2003; Coaker et al., 2005; Mackey et al., 2003), possible ADP-ribosylation by HopF2 (Wang et al., 2010; Wilton et al., 2010), and differential phosphorylation in the presence of AvrB and AvrRpm1 (this study). The proteolysis and phosphorylation events target an overlapping short domain on RIN4, the C-terminal NOI2 domain, which is part of a family of proteins cleaved by AvrRpt2 (Chisholm et al., 2005). *Arabidopsis* encodes ~15 paralogous NOI domain-containing proteins. Positions analogous to RIN4 T166 and F169 are nearly invariant within the NOI2 domains of 58 RIN4 orthologs (<http://www.phytozome.org/>; Cluster #23252144) and across 91 additional proteins orthologous to the remaining *Arabidopsis* NOI2-containing paralogs across the plant kingdom (<http://www.phytozome.org/>; Clusters #23252690, #23250407, and #23251786). Both AvrB and AvrRpm1 can promote virulence in plants lacking RIN4, indicating the existence of additional targets that may include other NOI-containing proteins (Belkhadir et al., 2004). Thus, we hypothesize that AvrB and AvrRpm1 suppress MTI by targeting RIN4 and additional NOI2-containing proteins and that T166 and F169, or equivalent residues, are central to these interactions. By extension, NOI2 domain-containing *Arabidopsis* proteins other than RIN4 also are likely to have roles in regulating plant defense. Our findings focus future experiments on this domain in RIN4 and its paralogues; the kinase(s) that phosphorylate RIN4 and, possibly, other NOI2 domain-containing proteins; and the precise mechanism(s) by which AvrB and AvrRpm1 modulate this phosphorylation event. Furthermore, the definition of functions for the other NOI2 domain family members are likely to contribute to understanding of plant immune system function.

EXPERIMENTAL PROCEDURES

Vectors

All cloning was performed using the Gateway system (Invitrogen, Carlsbad, CA). AvrB/AvrB G2A and AvrRpm1/AvrRpm1 ORFs were cloned with direct fusions of influenza hemagglutinin (HA) epitope tag at the C terminus into pDONR207 vector (Invitrogen). To generate estradiol-inducible constructs, each effector was cloned into the pMDC7 vector via an LR reaction. A T7 epitope tag (MASMTGGQMG) (Day et al., 2005) was added between the

RIN4 promoter (1.6 kb) and a genomic *RIN4* fragment (1.2 kb). The *RIN4* promoter was amplified with 5'- and 3'-primers that contain the T7 epitope tag as an overhang sequence. Gene-specific primers for genomic *RIN4* were generated to incorporate a T7 epitope sequence directly at the N terminus of genomic *RIN4* using the native stop codon in the 3'-primer. These full-length genomic *RIN4* constructs with the native promoter and T7 epitope tag were subcloned into pDONR207 vector. All AvrB binding site (BBS) mutants of RIN4 were generated by site-directed mutagenesis using wild-type genomic RIN4 as a template. To clone the genomic RIN4 construct into the binary vector, we generated a pBAR1-GW destination vector by inserting a Gateway cassette (Invitrogen) into the multicloning site of pBAR1, followed by restriction and ligation with HindIII and SacI fragments of pBAR1. Genomic *RPM1* driven by the native promoter in the pGPTV-HPT binary vector was used.

Plants

N. benthamiana for Agrobacterium-mediated transient assays were sown in soil (Day et al., 2005, 2006) and germinated in the greenhouse at 24°C with a long day photoperiod (16 hr light/8 hr dark). Two-week-old seedlings were transplanted to 4 inch square pots (one seedling per pot) and grown for 5–6 weeks before infiltration with Agrobacteria. For all transient assays, fully expanded leaves, which are the 3rd to the 5th from the first leaf at the bottom, were utilized.

Arabidopsis Col-0 wild-type and isogenic mutants were sown and grown as described (Boyce et al., 1998). To generate transgenic plants transformed with RIN4 BBS mutants, a *RPM1myc rpm1 rps2 rin4* line was generated by crossing the *RPM1-myc rpm1-3* (line AT5) (Boyce et al., 1998) with *rpm1 rps2 rin4* (Belkhadir et al., 2004) (see Supplemental Information).

Agrobacterium-Mediated Transient Assay in *Nicotiana benthamiana*

To reconstruct AvrB- or AvrRpm1-mediated, RPM1-dependent HR with RIN4 and RPM1 in *N. benthamiana*, a three-way *A. tumefaciens* infiltration was used. Strains of C58C1 (pCH32) transformed with AvrB or AvrRpm1 and their derivatives expressed in pMDC7 vector, the same strain carrying RIN4 and its derivatives in pBAR1GW binary vector, and RPM1 in pGPTV-HPT binary vector were infiltrated into the abaxial side of 5- to 6-week-old *N. benthamiana* leaves by hand infiltration with a 1 ml needleless syringe (see Supplemental Information).

Protein Extraction and Immunoblot Analyses

To examine the expression of proteins in transient assays and transgenic *Arabidopsis* plants, three leaf discs from infiltrated areas on *N. benthamiana* or two leaves of similar size from independent *Arabidopsis* transgenic lines were harvested and ground in liquid nitrogen. Total plant crude extracts were prepared with 150 μ l grinding buffer (20 mM Tris-HCl [pH 7.5], 150 mM NaCl, 1% Triton X-100, 1 mM EDTA [pH 8.0] and 0.1% SDS) with the addition of 10 mM DTT and 1 \times plant protease inhibitor cocktail (Sigma-Aldrich). The lysates were centrifuged at 14,000 rpm for 10 min at 4°C. Supernatants were collected and the concentration was determined with the BioRad Bradford quantification.

Protein extracts were electrophoresed through 8% and 13% SDS-PAGE, followed by transferring with semidry method (Thermo Scientific). Immunoblots were performed with a 1:5000 dilution of α -T7-HRP (Novagen), 1:2000 dilution of α -HA (Roche), and 1:10 dilution of α -myc (UNC Tissue Culture Facility). Blots were reacted with HRP-conjugated HRP secondary antibody and detected by ECL or ECL plus, following the manufacturer's guidance (GE Healthcare, Amersham).

Microsomal Fractionation and Two-Phase Partitioning

The microsomal fraction was extracted based on Boyce et al., 1998. For aqueous two-phase partitioning, the microsomal fraction was used to separate plasma membrane and endomembrane fraction as described previously (Kawasaki et al., 2005). Aqueous two-phase partitioning was done with a polymer concentration of 6.6% (wt/vol). α -ATPase (Agrisera) and α -BIP (Santa Cruz Biotechnology) antibodies were used as controls for plasma membrane and endomembrane fraction, respectively.

Bacterial Growth Assay in Planta

Pto DC3000(*avrB*) and (*avrRpm1*) were grown on KB media (10 g glycerin, 10 g peptone, 10 g tryptone, 10 ml 10% K₂HPO₄, 10 ml 10% MgSO₄, and 15 g agar

per 1 l) with appropriate antibiotics (100 µg/ml rifampicin and 25 µg/ml kanamycin) for 2 days. To measure the growth of *Pto* DC3000(EV), the same method was employed except that the amount of initial inoculum was 10⁴ cfu/ml. Statistical difference in bacterial growth at day 3 was analyzed by pairwise comparisons for all means using one-way ANOVA followed by Tukey-Kramer HSD with JMP 7.0 software (SAS Institute, Inc.).

Staining and Quantification of Hypersensitive Response In Planta

HR triggered by *Pto* DC3000(*avrB*) and *Pto* DC3000(*avrRpm1*) was visualized by trypan blue staining and quantified by conductivity measurement. Bacteria suspensions from *Pto* DC3000 possessing either *AvrB* or *AvrRpm1* were prepared as for the growth assay except that the final concentration for infiltration was 5 × 10⁷ cfu/ml. For staining with trypan blue, half of each leaf was inoculated to compare the infiltrated and noninfiltrated zone for HR. Approximately 20 leaves were stained. To measure the conductivity from infiltrated leaves, four leaf discs were collected and submerged into 6 ml of double-distilled water with three replicates per sample (n = 12) and then measured by conductivity meter (Orion, model 130) with indicated time points.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, Supplemental References, six figures, and one table and can be found with this article online at doi:10.1016/j.chom.2011.01.009.

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