

# *Arabidopsis* TAO1 is a TIR-NB-LRR protein that contributes to disease resistance induced by the *Pseudomonas syringae* effector AvrB

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The type III effector protein encoded by avirulence gene B (AvrB) is delivered into plant cells by pathogenic strains of *Pseudomonas syringae*. There, it localizes to the plasma membrane and triggers immunity mediated by the *Arabidopsis* coiled-coil (CC)-nucleotide binding (NB)-leucine-rich repeat (LRR) disease resistance protein RPM1. The sequence unrelated type III effector avirulence protein encoded by avirulence gene Rpm1 (AvrRpm1) also activates RPM1. AvrB contributes to virulence after delivery from *P. syringae* in leaves of susceptible soybean plants, and AvrRpm1 does the same in *Arabidopsis rpm1* plants. Conditional overexpression of AvrB in *rpm1* plants results in leaf chlorosis. In a genetic screen for mutants that lack AvrB-dependent chlorosis in an *rpm1* background, we isolated TAO1 (target of AvrB operation), which encodes a Toll-IL-1 receptor (TIR)-NB-LRR disease resistance protein. In *rpm1* plants, TAO1 function results in the expression of the pathogenesis-related protein 1 (PR-1) gene, suggestive of a defense response. In RPM1 plants, TAO1 contributes to disease resistance in response to *Pto* (*P. syringae* pathovars tomato) DC3000(*avrB*), but not against *Pto* DC3000(*avrRpm1*). The *tao1-5* mutant allele, a stop mutation in the LRR domain of TAO1, posttranscriptionally suppresses RPM1 accumulation. These data provide evidence of genetically separable disease resistance responses to AvrB and AvrRpm1 in *Arabidopsis*. AvrB activates both RPM1, a CC-NB-LRR protein, and TAO1, a TIR-NB-LRR protein. These NB-LRR proteins then act additively to generate a full disease resistance response to *P. syringae* expressing this type III effector.

Exposure to pathogens has driven the evolution of a sophisticated plant immune system that controls complex defense mechanisms for limiting infection (1). Plants express disease resistance proteins whose activation results in the restriction of pathogen growth and, typically, a localized cell death termed the hypersensitive response (HR). The most common class of disease resistance proteins consists of a central nucleotide binding (NB) site and a C-terminal leucine-rich repeat (LRR). NB-LRR proteins can be further subdivided based on the presence of a coiled-coil (CC) or a *Drosophila* Toll-IL-1 receptor (TIR) domain at the N terminus. In response to infection by Gram-negative bacterial phytopathogens, specific NB-LRR proteins are activated via recognition of a pathogen-encoded type III effector protein. This recognition can occur in an indirect manner, whereby type III effectors are perceived via their modification of host target proteins. Hence, NB-LRR proteins can survey the integrity of intracellular host proteins targeted by type III effectors acting as virulence factors. When a particular type III effector activates an NB-LRR protein, that type III effector is defined as an “avirulence protein.” This interaction is specific, in that a single allele of a NB-LRR gene is typically responsible for mediating resistance to a pathogen isolate expressing a particular type III effector (1).

One exception to this rule is provided by the *Pseudomonas syringae* type III effector proteins avirulence gene B (AvrB) and avirulence gene RPM1 (AvrRpm1); each can elicit disease resistance mediated by the *Arabidopsis* CC-NB-LRR protein

RPM1 (2, 3). After their delivery into host cells by the type III secretion pilus, AvrB and AvrRpm1 undergo N-terminal myristoylation that facilitates association with the plant plasma membrane (4). Here, these type III effectors associate with the membrane-bound host target protein, RIN4 (RPM1-interacting protein 4) (5). AvrB-RIN4 or AvrRpm1-RIN4 interactions are correlated with the phosphorylation of RIN4 and activation of RPM1 (6). Interestingly, in *Glycine max* (soybean) resistance to pathogen strains expressing AvrB or AvrRpm1 can be encoded by two different CC-NB-LRR genes, *Rpg1-b* and *Rpg1-r*, respectively (7, 8). Genetic data confirms that AvrB and AvrRpm1 have multiple host targets in *Arabidopsis rpm1* plants (9), suggesting that different NB-LRR receptors may have evolved to monitor the integrity of independent targets of these two type III effectors. Yet, the ability of AvrRpm1 and AvrB to trigger disease resistance in *Arabidopsis* has never been separated.

Transgenic *in planta* conditional expression of AvrB in *rpm1* plants leads to leaf chlorosis (4). AvrB-induced chlorosis is RIN4-independent (9), but requires the host protein RAR1 (required for Mla-resistance 1) (10). RAR1 is part of a chaperone complex that functions to positively regulate steady-state accumulation of numerous NB-LRR proteins (9, 11–13). Additionally, RAR1 is a positive regulator of basal disease resistance against virulent pathogens (13). Because AvrB can coimmunoprecipitate RAR1, the suggestion has recently arisen that RAR1 might be a virulence target of AvrB (10).

We performed a genetic screen to isolate mutants compromised for AvrB-induced chlorosis. This screen was performed in Mt-0, an ecotype (inbred genetic line) of *Arabidopsis* from which RPM1 is naturally deleted (14), but which responds to AvrB with leaf chlorosis (4). We defined and isolated TAO1 (target of AvrB operation), a gene encoding a disease resistance protein of the TIR-NB-LRR class. TAO1 activation correlates with defense gene expression. A specific mutant allele of TAO1 posttranscriptionally suppresses the steady-state accumulation of RPM1. In the Col-0 (RPM1) accession, TAO1 is required for a full resistance response to *P. syringae* pathovars tomato (*Pto*) DC3000(*avrB*) but not to *Pto* DC3000(*avrRpm1*), showing that recognition of these two type III effectors is genetically separable in *Arabidopsis*. TAO1-dependent phenotypes, like disease

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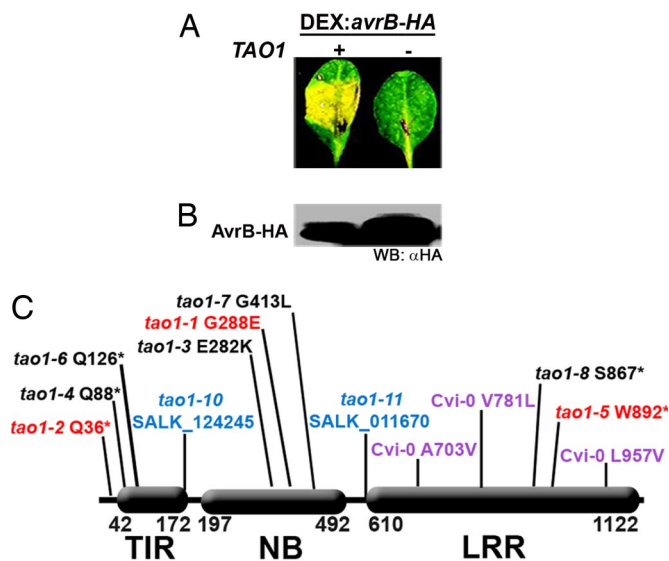
Data deposition: The sequences reported in this paper have been deposited in the GenBank database (accession nos. EU031442 and EU031443).

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**Fig. 1.** TAO1 is a TIR-NB-LRR R protein required for AvrB-induced chlorosis in *rpm1* host plants. (A) Mt-0 leaves were inoculated with *Agrobacterium* containing T-DNA with a *DEX:avrB-HA* transgene (4). Leaves were treated with DEX 48 h after inoculation. The picture was taken 72 h after inoculation. (B) Western blot showing accumulation of AvrB-HA at 8 h after DEX treatment. (C) Deduced structure of TAO1 alleles recovered in Mt-0 (red and black), Col-0 Salk T-DNA insertion lines (blue), and as polymorphisms in Cvi-0 (purple). For all missense mutations, the wild-type Mt-0 residue is listed first. TIR, amino acids 42–172. NB, amino acids 197–492. LRR, amino acids 610–1122. All TAO1 alleles in Mt-0 were generated by EMS mutagenesis except *tao1-8*, which is a fast neutron deletion of 1 bp in codon S667. Red *tao1* alleles represent alleles that were out-crossed away from the *DEX:avrB-HA* transgene and used for further analyses. The *tao1-10* (Salk\_124245) insertion begins at amino acid 168. The *tao1-11* (Salk\_011670) insertion begins at amino acid 597. The only amino acid difference between Mt-0 and Col-0 is V489M. Genomic TAO1 sequences for Mt-0 (EU031442) and Cvi-0 (EU031443) have been deposited in GenBank.

resistance mediated by all TIR-NB-LRR proteins, requires phytoalexin deficient 4 (PAD4). Our data provide further evidence that CC and TIR subclasses of NB-LRR proteins can function additively in disease resistance responses against biotrophic pathogens.

## Results

**TAO1 Encodes a TIR-NB-LRR Protein Required for AvrB-Induced Chlorosis in *rpm1* Host Plants.** We designed a conditional genetic screen to isolate loci required for AvrB-induced chlorosis in Mt-0 (*rpm1*). A stable single insertion transgenic line carrying a dexamethasone (DEX)-inducible *avrB* expression system [*DEX:avrB-HA* (15)] was established in Mt-0 (see *Methods*). Seed from this line was mutagenized, and  $\approx 400,000$  M2 plants were screened for loss of AvrB-induced chlorosis after DEX treatment (Fig. 1A). DEX-inducible AvrB-HA expression was confirmed in mutant (fully green) M3 progeny by Western blot analysis (Fig. 1B). We used transient *Agrobacterium* transformation of *DEX:avrB-HA* on several M3 progeny from each putative M2 mutant plant to eliminate mutations in the *DEX:avrB-HA* expression system. In this assay, plants mutated in the *DEX:avrB-HA* transgene were complemented by the transient expression of *DEX:avrB-HA* and display chlorosis after DEX treatment. These lines were not studied further.

Genetic complementation crosses indicated eight allelic, recessive *tao1* mutants that were also allelic to the naturally occurring *tao1* phenotype of Cvi-0 (4). Transgenic *DEX:avrB-HA rpm1 tao1-2* (Mt-0) and *DEX:avrB-HA rpm1 tao1-5* (Mt-0) lines were crossed independently to *rpm1-3 TAO1* (Col-0) for map-

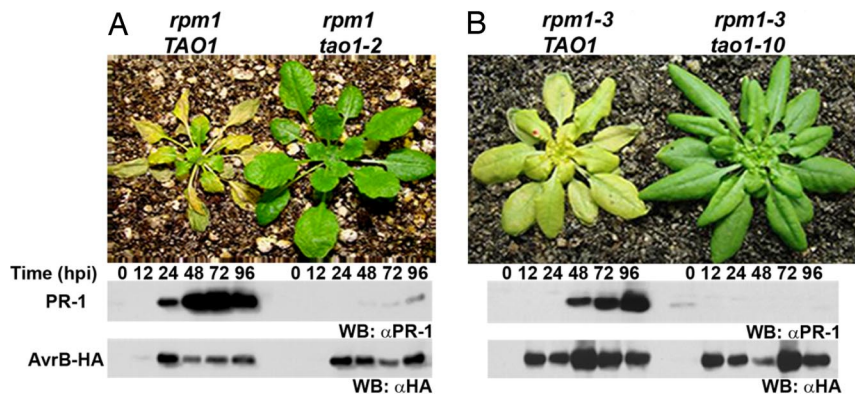
based cloning of TAO1. After DEX induction, F<sub>1</sub> plants exhibited host cell chlorosis, confirming that both *tao1-2* and *tao1-5* were recessive (data not shown). We isolated nonchlorotic plants in the F<sub>2</sub> generation. Two genotypes were present among these nonchlorotic F<sub>2</sub> plants: (i) those that lacked the *DEX:avrB-HA* transgene and (ii) those that contained the *DEX:avrB-HA* transgene and were homozygous *tao1*. We performed PCR with *avrB*-specific primers to distinguish between these classes; plants lacking the *DEX:avrB-HA* transgene were discarded. From the *tao1-5* cross,  $\approx 1,000$  informative F<sub>2</sub> plants showed that TAO1 was tightly linked to the bottom of chromosome 5. In a smaller F<sub>2</sub> population, the *tao1-2* mutant allele also mapped to the same location, which was in agreement with data previously presented for the Cvi-0 loss of function allele of TAO1 (4). To avoid mis-scoring caused by transgene silencing, we used transient assays with *Agrobacterium tumefaciens* (*DEX:avrB-HA*) of F<sub>3</sub> progeny to confirm the *tao1* genotypes of F<sub>2</sub> individuals. This approach, combined with the sequence identification of mutations in eight *tao1* alleles (designated *tao1-1* to *tao1-8*; Fig. 1C), identified TAO1 as the Mt-0 allele of the Col-0 gene At5g44510.

TAO1 encodes a TIR-NB-LRR protein of the TNL-G clade (16). There are no other NB-LRR genes at the TAO1 locus in the reference Col-0 genome sequence. The deduced TAO1 protein is only 26% similar to RPM1 and 25% similar to RPG1-B (soybean). For further analyses, we chose three alleles: *tao1-2*, an early stop mutation after Q36 and a likely null; *tao1-1*, a missense mutation at G238E in the NB domain; and *tao1-5*, a nonsense mutation at W892 that results in the deletion of the last five LRRs (13 through 17) (Fig. 1C). Sequencing of the naturally occurring *tao1* allele in Cvi-0 revealed three amino acid changes compared with the Mt-0 TAO1 sequence at 703, 781, and 957 (Fig. 1C). Col-0 TAO1 shares these amino acids with Mt-0, while also having a polymorphism at amino acid 489.

**TAO1 Activation Results in Expression of Pathogenesis-Related Protein-1 (PR-1).** We next evaluated whether TAO1 activation resulted in expression of a common defense response gene, PR-1. Five days after induction with DEX, *DEX:avrB-HA rpm1 tao1-2* plants were nonchlorotic, whereas the *DEX:avrB-HA rpm1 tao1-2* plants displayed strong leaf chlorosis (Fig. 2A). TAO1-mediated chlorosis correlated with an increase in PR-1 expression in leaf tissue (Fig. 2A). Thus, TAO1 function results in the induction of a common defense response marker. Similar results were found for *DEX:avrB-HA rpm1 tao1-5* (data not shown).

To analyze TAO1 function in Col-0, a better-studied genetic background for disease resistance studies than Mt-0, we used two mutant alleles designated *tao1-10* (SALK\_124245), and *tao1-11* (SALK\_011670) that carry exon insertions in At5g44510 (Fig. 1D). Both are TAO1 transcript nulls [supporting information (SI) Fig. S1]. These lines were crossed to an isogenic *DEX:avrB-HA rpm1-3 TAO1* line, and *DEX:avrB-HA rpm1-3 tao1-10* and *DEX:avrB-HA rpm1-3 tao1-11* lines were selected from the resulting progeny. Five days after induction with DEX, the *DEX:avrB-HA rpm1-3 TAO1* plants were chlorotic, whereas the *DEX:avrB-HA rpm1-3 tao1-10* lines were nonresponsive (Fig. 2B). Furthermore, TAO1 activation correlated with an increase in PR-1 expression (Fig. 2B). Similar results were found for *DEX:avrB-HA rpm1-3 tao1-11* (data not shown). Hence, *tao1-10* and *tao1-11* lose TAO1 function in response to AvrB expression *in planta*.

**RPM1 Function Is Lost Specifically in *tao1-5*.** We next addressed whether TAO1 contributes to RPM1-mediated disease resistance. For this, we introduced an RPM1 transgene containing a C-terminal myc epitope tag, expressed from the native RPM1 promoter (17), into Mt-0 (*rpm1 TAO1*). A line carrying a single copy of the *RPM1-myc* transgene was selected and then crossed



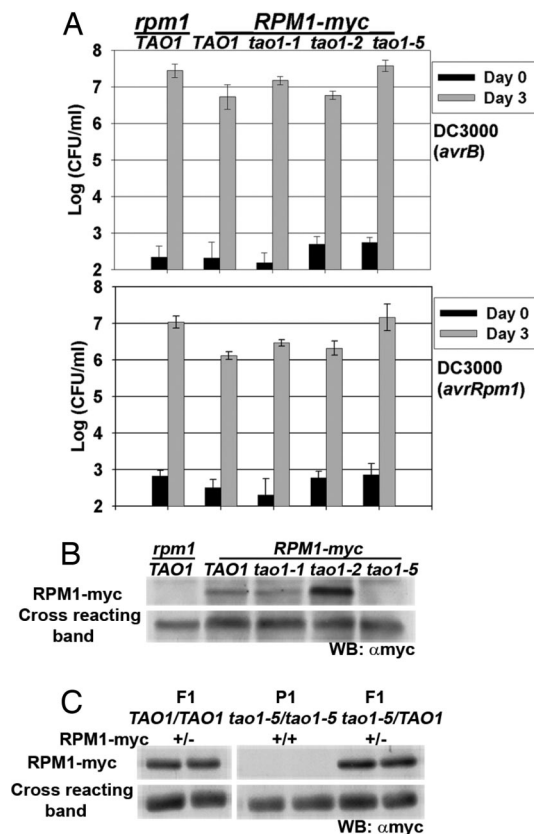
**Fig. 2.** TAO1 activation correlates with defense gene expression in both Mt-0 and Col-0. Five-week-old plants were sprayed with 20  $\mu$ M Dex to induce expression of AvrB-HA. (A) Mt-0 and *tao1-2*. (B) Col-0(*rpm1-3*) and *rpm1-3 tao1-10*. (Top) Photograph was taken 5 days after DEX treatment. (Middle) Western blot showing PR-1 protein levels. (Bottom) Western blot analysis showing AvrB-HA protein levels. Equivalent amounts of total proteins were loaded in all lanes (data not shown). The experiment is representative of three independent replicates.

into the *tao1-1*, *tao1-2*, and *tao1-5* mutant backgrounds (previously back-crossed and selected for loss of the *DEX:avrB-HA* transgene) (see *Methods*). Stable *RPM1-myc TAO1*, *RPM1-myc tao1-1*, *RPM1-myc tao1-2*, and *RPM1-myc tao1-5* lines were selected, and RPM1 function was assessed. After hand infiltration with *Pto* DC3000(*avrB*) or *Pto* DC3000(*avrRpm1*), RPM1 function resulted in only a 5- and 10-fold growth restriction of bacteria, respectively (Fig. 3A), which is much less bacterial growth restriction than observed in Col-0 (*RPM1*) (Fig. 4A). This weak RPM1-mediated growth restriction is likely caused by lower accumulation of RPM1-myc in Mt-0 relative to Col-0 (Fig. S2). Lower RPM1-myc accumulation may be the result of genetic background differences between Mt-0 and Col-0 that control RPM1 accumulation or transgene positional effects. In *RPM1-myc tao1-1* and *RPM1-myc tao1-2*, there was essentially no effect on the weak *RPM1*-mediated growth restriction of *Pto* DC3000(*avrRpm1*) or *Pto* DC3000(*avrB*) (Fig. 3A). Interestingly, RPM1 function was lost in the *RPM1-myc tao1-5* background in response to either *Pto* DC3000(*avrB*) or *Pto* DC3000(*avrRpm1*) (Fig. 3A).

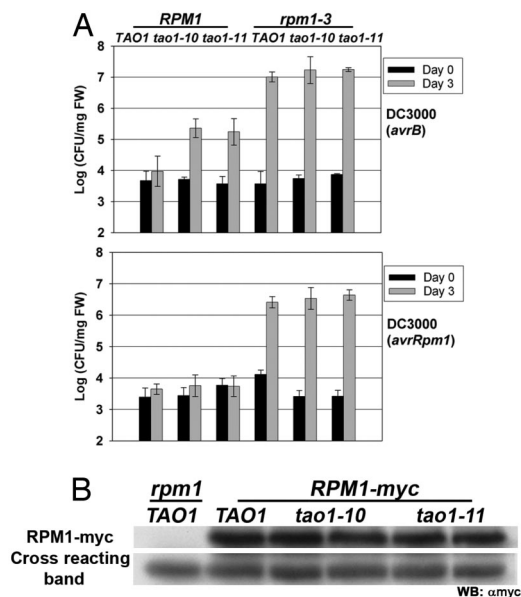
**RPM1 Does Not Accumulate in the *tao1-5* Background.** We postulated two possible reasons why RPM1 function is lost in the *tao1-5* background. The *tao1-5* mutant protein could either prevent RPM1 signaling, or the *tao1-5* mutant protein might negatively regulate RPM1 steady-state accumulation. To distinguish between these possibilities, the *RPM1-myc TAO1*, *RPM1-myc tao1-1*, *RPM1-myc tao1-2*, and *RPM1-myc tao1-5* lines were subjected to Western blot analysis for RPM1-myc. Despite some experimental variation, RPM1 accumulated normally in *RPM1-myc tao1-1* and *RPM1-myc tao1-2* (Fig. 3B). By contrast, RPM1 accumulation was abolished in *RPM1-myc tao1-5* (Fig. 3B). RT-PCR analysis revealed that this effect on RPM1 is posttranscriptional (Fig. S3). Interestingly, the effect of *tao1-5* on RPM1 accumulation was recessive (Fig. 3C).

**TAO1 Is Required for Full Disease Resistance Against *Pto* DC3000(*avrB*) in Col-0.** We introgressed the *tao1-10* and *tao1-11* null alleles into *rpm1-3*. Homozygous *rpm1-3 tao1-10* and *rpm1-3 tao1-11* lines were selected to evaluate TAO1 function in both the presence and absence of RPM1. Interestingly, after dip inoculation in a bacterial suspension of *Pto* DC3000(*avrB*), both the *RPM1 tao1-10* and *RPM1 tao1-11* lines displayed  $\approx$ 10-fold less disease resistance than *RPM1 TAO1* (Fig. 4A). This effect of TAO1 was not seen with hand infiltrations (data not shown), suggesting that TAO1 may act early in pathogen infection. Hence, TAO1 contributes  $\approx$ 1–5% of the full resistance to *Pto* DC3000(*avrB*) in

the presence of RPM1. We observed no enhanced susceptibility to *Pto* DC3000(*avrB*) in the *rpm1-3 tao1-10* and *rpm1-3 tao1-11* lines, indicating that the weak TAO1 contribution to disease



**Fig. 3.** Weak RPM1 function in Mt-0 is suppressed posttranscriptionally by *tao1-5*. (A) Four-week old plants were hand-infiltrated with *Pto* DC3000(*avrB*) (Upper) or *Pto* DC3000(*avrRpm1*) (Lower) at  $10^5$  cfu/ml. Error bars represent the SD among four samples. This experiment is indicative of three independent replicates. Similar results were seen with dip infections (data not shown). (B and C) Western blot analysis showing RPM1-myc accumulation in plant lines with the noted genotypes. Cross-reacting band indicates equivalent amounts of total protein loading. Samples from hybrid (F<sub>1</sub>) and parental (P<sub>1</sub>) lines are noted above their respective genotypes. These experiments are indicative of four (B) and two (C) independent replicates. The higher than wild-type level of RPM1-myc accumulation in the *tao1-2* background shown in B was not reproducible.



**Fig. 4.** TAO1 contributes to disease resistance against *Pto* DC3000(*avrB*) but not *Pto* DC3000(*avrRpm1*). (A) Two- to three-week-old plants were dip-infiltrated with *Pto* DC3000(*avrB*) (Upper) or *Pto* DC3000(*avrRpm1*) (Lower) at a bacterial concentration of  $2.5 \times 10^7$  cfu/ml. Error bars represent the SD among four samples. This experiment is representative of three independent replicates. (B) Western blot analysis showing RPM1-myc accumulation in plant lines with the noted genotypes. The cross-reacting band serves as an indicator of equivalent amounts of total protein loaded. This experiment is representative of two independent replicates.

resistance is too low to measure in the absence of RPM1 (Fig. 4A). We observed full RPM1-mediated growth restriction in the *RPM1 tao1-10* and *RPM1 tao1-11* lines after dip inoculation of *Pto* DC3000(*avrRpm1*) (Fig. 4B). Hence, TAO1 contributes specifically to disease resistance triggered by *Pto* DC3000(*avrB*) and does not contribute to RPM1-mediated disease resistance in response to *Pto* DC3000(*avrRpm1*). As with *Pto* DC3000(*avrB*), we observed no enhanced susceptibility of the *rpm1-3 tao1-10* and *rpm1-3 tao1-11* lines to *Pto* DC3000(*avrRpm1*) (Fig. 4A).

Although RPM1-mediated resistance to *Pto* DC3000(*avrRpm1*) was unaffected in the *tao1-10* and *tao1-11* lines, the possibility existed that TAO1 affects RPM1 steady-state accumulation. This scenario would indicate that different accumulation levels of RPM1 are required for resistance to *Pto* DC3000(*avrB*) versus *Pto* DC3000(*avrRpm1*). To address this possibility, the *rpm1-3 tao1-10* and *rpm1-3 tao1-11* lines were crossed to a *rpm1-3 RPM1-myc TAO1* line (17). Stable *rpm1-3 RPM1-myc tao1-10* and *rpm1-3 RPM1-myc tao1-11* lines were selected. Western blot analysis revealed that RPM1 accumulation was equivalent in the *tao1-10* and *tao1-11* lines relative to the *TAO1* parental line (Fig. 4B). Hence, loss of TAO1 does not affect RPM1 steady-state accumulation. Therefore, the contribution of TAO1 to disease resistance against *Pto* DC3000(*avrB*) in Col-0 is likely to be at the signaling level.

**TAO1 Requires PAD4 for Function.** The *Arabidopsis* PAD4 protein functions in both basal defense and TIR-NB-LRR-mediated disease resistance (18, 19). To assess whether PAD4 is required for TAO1 function, we crossed the *DEX:avrB-HA rpm1-3* line to the *pad4-1* mutant line (20) and selected homozygous *F2 rpm1-3 pad4-1* lines that contained the *DEX:avrB-HA* transgene. After inducing AvrB expression *in planta*, we observed that both TAO1-mediated chlorosis and PR-1 expression were lost in the *rpm1-3 pad4-1* lines (Fig. S4A). Furthermore, inoculation with *Pto* DC3000(*avrB*) and *Pto* DC3000(*avrRpm1*) on *pad4-1 TAO1*

*RPM1* lines revealed that the TAO1 defense function against *Pto* DC3000(*avrB*) is also lost in a *pad4-1* mutant (Fig. S4B). Thus, these results demonstrate that TAO1 requires PAD4 for function.

## Discussion

AvrB is a bacterial type III effector protein that causes modification of the *Arabidopsis* host protein RIN4, subsequently activating the NB-LRR protein RPM1 (5, 6, 21) or contributing to pathogen virulence in susceptible plants (7). The sequence-independent type III effector AvrRpm1 acts similarly (22). In the absence of RPM1, *in planta* expression of AvrB produces a host cell chlorosis originally speculated to be indicative of an AvrB virulence function (4). Because AvrB-mediated chlorosis is RIN4 independent, AvrB must have additional targets within the *Arabidopsis* cell (9). Our aims were to (i) identify host proteins required for AvrB-induced chlorosis, (ii) assess whether and how these host components affect RPM1 function triggered by either AvrB or AvrRpm1, and (iii) clarify whether these host factors are specific for the responses to AvrB. We identified only one gene in our mutant screen, and it encodes TAO1, a TIR-NB-LRR disease resistance protein.

Our data demonstrate that TAO1 activation contributes to disease resistance in response to *Pto* DC3000(*avrB*), but not in response to *Pto* DC3000(*avrRpm1*) (Fig. 4A). To explain the functional relationship between AvrB, TAO1, and RPM1, we propose the following model (Fig. S5). In *rpm1* plants, TAO1 activation in response to AvrB is insufficient to cause a HR (data not shown) or significantly restrict *Pto* DC3000(*avrB*) growth (Fig. 4A). However, chlorosis and expression of a common defense protein, PR-1, are induced (Fig. 2; see also Fig. S5 Left). These data demonstrate that TAO1 is a weak disease resistance protein that cannot create sufficient defense response amplitude to restrict *Pto* DC3000(*avrB*) growth in the absence of RPM1. In *tao1* plants, RPM1 activation in response to AvrB results in a HR in addition to a significant restriction of *Pto* DC3000(*avrB*) growth (Fig. 4A) (Fig. S5 Center). Finally, the additive functions of RPM1 and TAO1 facilitate full disease resistance to *Pto* DC3000(*avrB*) (Fig. 4A) (Fig. S5 Right).

Because TAO1 function is RIN4 independent (9) and RIN4 is required for RPM1 function (5), TAO1 perception of AvrB might occur at a cellular site that lacks RIN4 and RPM1. Recently, two studies (6, 21) identified AvrB mutants that are unable to activate RPM1, but still activate TAO1. For example, the AvrB T125A mutation abolished both the RIN4-AvrB interaction and the ability to trigger RPM1, but retained the ability to activate TAO1 (6, 21). Additionally, the AvrB D297A mutation retained interaction with RIN4 and lost RPM1 activation but still elicited TAO1 (6, 21). These data suggest that there are likely to be structural differences in how AvrB targets host proteins and consequently triggers either RPM1 or TAO1. Despite the potentially separate mechanisms of TAO1 and RPM1 activation, both require AvrB membrane localization via myristoylation (4). These data may indicate that like RPM1 and RIN4, a second putative protein(s), targeted by AvrB and associated with TAO1, is also present at the plasma membrane.

Interestingly, the steady-state levels of RPM1 are significantly diminished in *tao1-5* (Fig. 3B). Because the other loss-of-function mutant alleles tested, *tao1-1* (presumptive P-loop dead) and *tao1-2* (presumptive null), do not affect RPM1 accumulation and *tao1-5* homozygosity is required for its effects on RPM1, *tao1-5* acts as a recessive gain-of-function allele (Fig. 3 B and C). This phenotype is reminiscent of the effects of specific alleles of *hsp90.2* on RPM1 accumulation (23, 24). The *tao1-5* allele exhibits no other observable phenotypic defects (data not shown). A *tao1-5*-encoded protein could sequester RPM1 directly or negatively affect RPM1 stability through its effects on an intermediate host protein. Three host proteins shown to be

required for RPM1 steady-state accumulation are RIN4 and the general NB-LRR chaperone complex members RAR1 and HSP90.2 (5, 12). Because neither RPS2 function nor RPS5 steady-state accumulation are compromised in the *tao1-5* background (data not shown), it is unlikely that the *tao1-5* mutant protein sequesters RIN4 or RAR1. Hence, we speculate that HSP90.2 or an as-yet-undiscovered protein that contributes to RPM1 steady-state accumulation is altered by the mutant *tao1-5*-encoded product.

Recently, Shang *et al.* (10) demonstrated that RAR1 is required for AvrB-dependent chlorosis in *rpm1* plants (10). They also showed that AvrB-induced chlorosis correlated with increased growth of a *Pto* DC3000(*hrpL*) mutant that lacks the type III pilus and therefore cannot deliver type III effectors to the plant cell. This relaxation of basal defense responses was RAR1 dependent. Also, when overexpressed, AvrB and RAR1 can be coimmunoprecipitated (10) and also allow reactivation of a split-luciferase reporter (albeit under conditions where the AvrB myristoylation site is buried in the reporter fusion and mislocalization of AvrB is likely) (25). These results led the authors to conclude that RAR1 is a “virulence target” of AvrB, although no direct interaction of AvrB and RAR1 was demonstrated (10). Our data demonstrate that AvrB-induced chlorosis is mediated by the TIR-NB-LRR protein TAO1 and that TAO1 functions as a “weak *R* gene” (Figs. 2 and 4A). Thus, there is a contradiction: is the observed host cell chlorosis indicative of AvrB virulence targeting of RAR1, or does it represent a TAO1-mediated disease resistance response, which in turn requires RAR1?

There are several possible explanations for the differing interpretations presented in ref. 10 and here regarding what AvrB-mediated chlorosis represents. First, the increase in *Pto* DC3000(*hrpL*) growth observed after induced AvrB overexpression *in planta* was COI1 dependent, whereas AvrB-induced chlorosis was COI1 independent (10). Hence, our observation of AvrB-mediated, TAO1-dependent chlorosis can be genetically separated from AvrB-mediated, COI1-dependent suppression of basal defense (10). Second, our data show that AvrB-mediated chlorosis requires the host protein PAD4 (Fig. S4), a positive regulator of both basal defense and TIR-NB-LRR-mediated disease resistance (18, 19). Third, a very strong regimen of conditional overexpression of AvrB [30  $\mu$ M DEX spray 2 days before the infection, and also continually throughout the *Pto* DC3000(*hrpL*) infection] was used in ref. 10. Using a regimen of two times 10  $\mu$ M DEX treatment, but only 1 day before infection, we were able to fully induce AvrB-dependent chlorosis in TAO1 plants, but did not observe increased growth of *Pto* DC3000(*hrcC*) (data not shown). These data suggest that the increased growth of *Pto* DC3000(*hrpL*) shown in ref. 10 is caused by extraordinary levels of AvrB expression *in planta*, resulting in extensive tissue collapse. Our data demonstrate that TAO1 contributes to the restriction of pathogen growth when AvrB is delivered naturally via a type III pilus from *Pto* DC3000 (Fig. 4A). Furthermore, we note that using the same experimental conditions as in ref. 10, it was shown that AvrB-mediated chlorosis correlated with a growth reduction of *Pto* DC3000 (33). This growth reduction was RAR1 dependent, consistent with a role for NB-LRR proteins in basal defense (13), but difficult to reconcile with the data presented in ref. 10. RAR1 regulates the function of many NB-LRR proteins by acting as a positive regulator of NB-LRR stability (9, 11–13). Hence, the simplest scenario to explain the sum of the data presented here and in ref. 10 is that the loss of AvrB-dependent chlorosis in a *rar1* background indicates that TAO1 requires RAR1 for steady-state accumulation and therefore function.

The bacterial type III effector proteins AvrB and AvrRpm1 each trigger strong disease resistance through RPM1. Yet the activation of TAO1 by AvrB suggests that additional layers of

recognition in response to infection can effectively add to the plant's defense response. In a mechanistic sense, if a given type III effector has multiple cellular targets, then direct or indirect recognition by more than one NB-LRR may add to both the flexibility and the amplitude of the host's overall response. These scenarios provide hosts with greater evolutionary capacity, by allowing NB-LRR proteins to continually fine-tune their responses to a single pathogen effector while maintaining an intermediate level of disease resistance. This possibility may explain recent examples of genetic requirements for combined action of (i) a TIR-NB-LRR and a CC-NB-LRR protein (26) and (ii) a TIR-NB-LRR and an atypical CC-containing NB-LRR (27) against a virus and an isolate of the oomycete parasite *Hyaloperonospora parasitica*, respectively. Our findings further the concept that a CC-NB-LRR and a TIR-NB-LRR, which typically trigger different response pathways in plants (28), can additively contribute to disease resistance.

## Methods

**A. tumefaciens-Mediated, DEX-Inducible Transient Expression Assays.** Transient transformation assays were performed as described (4). *Arabidopsis* chlorosis response in this assay was observed at 72 h after induction. Protein was extracted 8 h after induction from four transformed leaf discs.

**Generation of Stable Transgenic Lines, Mutagenesis, and Screening.** Stable transgenic plant lines were generated by the floral dip method (29). Segregation of hygromycin resistance was used to isolate lines containing single-copy insertions of both the *DEX:avrB-HA* (4) and *RPM1-myc* (17) transgenes in Mt-0. For ethane methyl sulfonate (EMS) mutagenesis,  $\approx 30,000$  seeds were incubated in a 0.25% EMS solution for 8 h, washed in distilled water, and dried on filter paper. Fast neutron mutagenesis was performed by the International Atomic Energy Agency research laboratory in Vienna, Austria. Approximately 700,000 M2 seeds in 100 lots (80 EMS, 20 fast neutron) were sprayed with 20  $\mu$ M DEX as described (30). Putative mutants were isolated and allowed to self-pollinate. M3 progeny were retested by *Agrobacterium* transient delivery of *DEX:avrB-HA* as described above.

**Positional Cloning of TAO1 and Out-Crossing of the DEX:avrB-HA Transgene.** Mapping of TAO1 was performed as described (31). Candidate genes within the TAO1 interval were PCR-amplified from the *tao1* mutants, and PCR products were sequenced directly. To obtain *tao1* lines lacking the *DEX:avrB-HA* transgene, M3 *tao1* lines were crossed to nontransgenic TAO1 Mt-0 plants. F<sub>2</sub> plants containing the *DEX:avrB-HA* transgene were selected and sprayed with 20  $\mu$ M DEX. Nonchlorotic (*tao1*) lines were selected and allowed to self-pollinate. F<sub>3</sub> individuals that lacked the *DEX:avrB-HA* transgene were selected and allowed to self-pollinate. F<sub>4</sub> seed was confirmed for *tao1* genotype by *Agrobacterium* transient transformation of *DEX:avrB-HA*. Subsequent confirmation of *tao1* mutant alleles was performed by using dominant cleaved amplified polymorphic sequences specific to the *tao1* allele in question.

**Bacterial Infections.** For Fig. 3A, 4-week-old plants were hand-infiltrated with bacteria at a concentration of  $10^5$  cfu/ml. For each sample, four leaf disks were pooled per time point (16 total). Leaf disks were ground in 10 mM MgCl<sub>2</sub>, subject to serial dilutions, and plated on King's Broth (KB) plates with rifampicin selection. For Fig. 4A and Fig. S4B, dip inoculations were performed as described (32). Briefly, 2- to 3-week-old seedlings were inoculated by dipping in solutions of bacteria at a concentration of  $2.5 \times 10^7$  cfu/ml. Three seedlings were placed in 1 ml of 10 mM MgCl<sub>2</sub> with 0.2% Silwet L-77. The weights of the tubes with seedlings were measured, and the tubes were then shaken for 1 h at 28°C. Serial dilutions in 10 mM MgCl<sub>2</sub> were made and plated on KB plates with rifampicin selection.

**Protein Analysis.** For detection of AvrB-HA, PR-1, and RPM1-myc, total protein extracts were prepared by grinding  $\approx 200$  mg of leaf tissue in buffer [50 mM Tris-HCl (pH 8.0), 1% SDS, 1 mM EDTA, 1 mM 2-mercaptoethanol, and plant protease inhibitor mixture (Sigma)]. Protein concentrations were determined by using the Bradford-Lowry method and quantification buffer from BioRad. Ten microliters of  $6 \times$  Laemmli buffer (final concentration  $1 \times$ ) was added to all measured samples after quantification. For immunodetection of AvrB-HA and PR-1, 10- $\mu$ g protein samples were electrophoresed on 14% SDS polyacrylamide gels. For immunodetection of RPM1-myc, 40- $\mu$ g protein samples were electrophoresed on 8% SDS polyacrylamide gels. Western blots were per-

formed by using standard methods. Anti-PR-1 serum (gift of Robert A. Dietrich, Syngenta, Research Triangle Park, NC) was used at a dilution of 1:10,000. The detection of the HA epitope tag was with supernatants from cultures of hybridoma 3F10 monoclonal anti-HA antibody (Roche) at a dilution of 1:1,000. Detection of the myc epitope tag was with supernatants from hybridoma 9E10 monoclonal anti-myc antibody at a dilution of 1:10 (17).

**RT-PCR.** RNA was isolated from various plants lines by using TRIZOL Reagent (GIBCO/BRL) according to the manufacturer's instructions. RT-PCR (RETROscript; Ambion) analysis in Fig. S1 and Fig. S3 was performed according to the manufacturer's instructions. Plant 185 Competimer primers (Ambion) were used to coamplify the 18S internal loading control. For Fig. S1, primers flanking the first

intron of *TAO1* 5'-tatgaatgcagagaagattgg-3' and 3'-ataaccgttctgctgtagag-5' were used with the PCR conditions 94°C 10 s, 55°C 60 s, 72°C 60 s, 31 cycles. A ratio of 9:1 Competimer/18S was used. For Fig. S3, the RPM1 transcript primers 5'-caccatgctcgctactgttatttgg-3' and 3'-cacttgcacgcatcaatagg-5' RPM1 transcript primers were used with the PCR conditions 94°C 10 s, 55°C 60 s, 72°C 60 s, 31 cycles. A ratio of 9.5:1 Competimer/18S was used. Products were separated on a 1% agarose gel.

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