Title: Pattern-recognition receptors are required for NLR-mediated plant immunity

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Abstract

The plant immune system is fundamental to the survival and productivity of plants in crop fields and natural ecosystems. Substantial evidence supports the prevailing notion that plants have evolved two branches of innate immune signaling, called pattern-triggered immunity (PTI) and effector-triggered immunity (ETI). PTI is triggered by conserved microbial patterns via surface-localized pattern-recognition receptors (PRRs), whereas ETI is activated by pathogen effector proteins via mostly intracellularly-localized receptors called nucleotide-binding, leucine-rich...
repeat proteins (NLRs)\(^1\). PTI and ETI have traditionally been considered to be separate immune signaling pathways and have evolved sequentially\(^5,6\). Here we show, contrary to the perception of PTI and ETI being separate immune signaling pathways, Arabidopsis PRR/co-receptor mutants, \(fls2/efr/cerkl\) and \(bak1/bkk1/cerkl\) triple mutants, are greatly compromised in ETI responses triggered by \(Pseudomonas syringae\) effectors. We further identified the NADPH oxidase (RBOHD)-mediated production of reactive oxygen species as a critical early signaling event connecting PRR and NLR signaling cascades and show that PRR-mediated activation of RBOHD is necessary for full activation of RBOHD during ETI. Furthermore, NLR signaling rapidly augments the transcript and protein levels of key PTI components at an early stage and in a salicylic acid-independent manner. Our study supports an alternative model in which PTI is in fact an indispensable component of ETI during bacterial infection, implying that ETI halts pathogen infection, in part, by directly co-opting the anti-pathogen mechanisms proposed for PTI. This alternative model conceptually unites two major immune signaling pathways in the plant kingdom and mechanistically explains the long-observed similarities in downstream defense outputs between PTI and ETI.

Main

It is generally believed that PRRs and NLRs, with largely distinct activation mechanisms and subcellular localizations, mediate different immune signaling pathways. PRRs are surface-localized receptor-like kinases/proteins (RLKs/RLPs) with extracellular ligand-binding domain to sense conserved molecular patterns, ranging from bacterial flagellin and fungal chitin molecules from both pathogenic and nonpathogenic microbes. NLRs, on the other hand, are intracellular proteins that sense pathogen-derived effector proteins inside the plant cell and can be further classified into the coiled coil (CC)-type or Toll/interleukin-1 receptor (TIR)-type, depending on their N-terminal domain\(^7\). However, PRR- and NLR-mediated signaling pathways result in many similar downstream immune outputs, including defense gene expression, production of reactive oxygen species (ROS) and callose deposition at the plant cell wall\(^8,9\). The underlying reason is not clear and mechanistic relationship between the two immune pathways remains enigmatic. Notably, while many PRR signaling components have been identified and anti-pathogen mechanisms described, the downstream signaling events in ETI and how ETI halts...
pathogen growth remain largely elusive, despite recent breakthroughs in the understanding of NLR protein structures and activities\textsuperscript{10-13}.

**Requirement of PRR/co-receptors for ETI**

Using *Arabidopsis thaliana*-*Pseudomonas syringae* pathosystem, we accidentally discovered a striking and unexpected role of PRR/co-receptors in ETI. Specifically, an “avirulent”, ETI-eliciting bacterial strain, *P. syringae* pv. *tomato* (*Pst*) DC3000(*avrRpt2*), which activates RPS2 (Resistance to *P. syringae* 2)-dependent ETI in wild-type Col-0 plants\textsuperscript{14,15}, failed to elicit effective ETI in two separate PRR/co-receptor Arabidopsis mutants, *fis2/efr/cerk1* (*fec*) and *bak1/bkk1/cerk1* (*bbc*) mutants, which are mutated in major PRR/co-receptors recognizing bacteria-associated molecular patterns\textsuperscript{16}. As shown in Fig. 1a, the *fec* and *bbc* mutants did not mount an effective ETI against *Pst* DC3000(*avrRpt2*). To determine whether a requirement of PRR/co-receptors for ETI is specific to *Pst* DC3000(*avrRpt2*) or is a more general phenomenon, we tested two other ETI-triggering “avirulent” effectors, AvrPphB and AvrRps4, which are recognized by RPS5\textsuperscript{17} and RPS4\textsuperscript{18}, respectively, in Arabidopsis Col-0 accession. We found that the compromised ETI phenotype in *fec* and *bbc* mutants held true for both AvrPphB and AvrRps4 (Extended Data Fig. 1), suggesting a potentially broad role of PRR/co-receptors in ETI pathways. We subsequently focused on AvrRpt2-triggered ETI for in-depth characterization.

Hypersensitive response (HR), manifested by fast cell death under high bacterial inoculum, is a hallmark of ETI. We tested HR phenotype in *fec* and *bbc* mutants in response to *Pst* DC3000(*avrRpt2*) and found that, even though HR cell death eventually occurs in these mutants, the rate was delayed, as shown by the compromised tissue collapse 7h after bacteria infiltration (Fig. 1b).

For the past several decades, classical studies of ETI triggered by *Pst* DC3000 carrying “avirulent” effector genes have been performed in the presence of all 36 endogenous effector genes in *Pst* DC3000. Because many of the endogenous effectors in *Pst* DC3000 are linked to interference of PTI and/or ETI via unclear mechanisms\textsuperscript{19,20}, it is not always easy to clearly interpret the relationship between PTI and ETI during infection by avirulent *Pst* DC3000 strains. We took advantage of the recent availability of *Pst* DC3000 strain D36E\textsuperscript{21}, in which all 36 effector genes are deleted and therefore is expected to activate only PTI, and D36E(*avrRpt2*)
strain, which delivers only AvrRpt2 and activates both PTI and RPS2-mediated ETI with no interference from any endogenous Pst DC3000 effectors. Although D36E is greatly reduced in virulence compared to Pst DC3000 (Fig. 1a), we could still observe a robust AvrRpt2-induced ETI in Col-0 plants, with strain D36E(avrRpt2) growing significantly less than strain D36E (Fig. 1c). We found that AvrRpt2-triggered ETI was no longer detectable in both fec and bbc mutants (Fig. 1c), demonstrating that the requirement of PRR/co-receptors for AvrRpt2-triggered ETI was not caused by some hidden interactions between endogenous Pst DC3000 effectors and Arabidopsis fec and bbc mutants.

**Dual role of RBOHD in ROS during PTI and ETI**

Previous studies have shown that AvrRpt2 cleaves the plant protein RIN4 (RPM1-interacting protein 4), leading to activation of the NLR protein RPS2. To understand why AvrRpt2-mediated ETI is lost in fec and bbc mutants, we examined the cleavage of the RIN4 protein by AvrRpt2. Results showed that D36E(avrRpt2)-induced RIN4 protein depletion was normal in the fec and bbc mutants compared to that in Col-0 plants (Fig. 1d). Gene expression analysis showed that RPS2 transcript level is also similar in all genotypes after bacteria inoculation (Extended Data Fig. 2). We then sought to assay downstream signaling events of AvrRpt2-triggered ETI and examined MAPK phosphorylation level, which has been reported to be stronger and more sustained after AvrRpt2 recognition by RPS2, compared to that induced during PTI. As shown in Fig. 1e, we observed strong ETI-associated MPK3/6 phosphorylation (i.e., at 4 or 8 h post inoculation) in Col-0 plants, which however remained intact in fec and bbc mutants.

Another important immune response associated with both PTI and ETI is production of ROS, including superoxide and hydrogen peroxide (H$_2$O$_2$), which have been proposed to act as defense molecules that kill pathogens and signaling molecules that further activate immune responses. Using luminol-horseradish peroxidase (HRP)-based method, we examined PTI- and ETI-associated ROS production in a transgenic avrRpt2 plant, in which avrRpt2 expression is driven by a dexamethasone (DEX)-inducible promoter. In this system, PTI and ETI activation can be initiated separately or in combination using PAMP (e.g., flg22, a 22-aa peptide derived from bacterial flagellin) and DEX treatments in the absence of bacterial infection. As shown in Fig. 1f, flg22 alone triggered a fast and transient ROS burst as reported before, while DEX-induced
expression of AvrRpt2 alone triggered only a weak and kinetically slower ROS burst. Interestingly, co-treatment of flg22 and DEX triggered a strong and sustained second-phase ROS burst, peaking at 2h to 3h after treatment, and lasted for several hours (Fig. 1f, g), a profile that bears a remarkable similarity to previously observed during bacteria-triggered ETI. This result raises an intriguing possibility that activation of PTI may be required for the production of a strong and sustained ROS characteristic of ETI. To test this hypothesis, we generated bbc/DEX::avrRpt2 plants by transforming the DEX::avrRpt2 construct into the bbc mutant plant (as well as Col-0 plant as control). Independent lines in which the avrRpt2 expression level was similar or even higher than that in Col-0/DEX::avrRpt2 plants were chosen (Extended Data Fig. 3) for further analysis. As shown in Fig. 1h, i, in the bbc/DEX::avrRpt2 plants, not only flg22-induced first-phase ROS is absent, but also the second-phase AvrRpt2-triggered ROS burst is almost completely abolished, clearly demonstrating a requirement of PRR/co-receptors for ETI-associated ROS production.

To examine whether PTI- and ETI-associated ROS bursts are produced at the same or different subcellular compartments, ROS production was monitored with the fluorescent dye H$_2$DCFDA, which can cross the plasma membrane of the plant cell and detect both apoplastic and cytosolic ROS. As shown in Fig. 2a, strong fluorescent signal was detected in the apoplastic space of Col-0 leaves 5h post infiltration of D36E(avrRpt2). The apoplastic signal was much weaker in the bbc mutant plant, which was indistinguishable compared to the rps2 control plant infiltrated with D36E(avrRpt2) or Col-0 plant infiltrated with D36E (Fig. 2a). Two classes of enzymes, the NADPH oxidases such as respiratory burst oxidase homolog D (RBOHD) and peroxidases, have been shown to be involved in generating apoplastic ROS in pathogen-infected plant leaves. We therefore investigated which class is involved in generating AvrRpt2-triggered ROS by using chemical inhibitors diphenylene iodonium (DPI), which inhibits NADPH oxidases, and salicylhydroxamic acid (SHAM) and sodium azide, which inhibit peroxidase activities. As shown in Extended Data Fig. 4a-c, co-treatment of DPI, but not SHAM or sodium azide, with flg22 and DEX almost completely blocked ETI-associated ROS and greatly compromised PTI-associated ROS. Interestingly, when we added these inhibitors at 40 min after flg22+DEX treatment (i.e., after PTI-associated ROS and before the start of ETI-associated ROS), still only DPI, but not SHAM or sodium azide, almost completely blocked ETI-ROS (Fig. 2b), indicating...
that NADPH oxidases mediate ETI-associated ROS. We further tested whether NADPH oxidase RBOHD, which has been shown to play a prominent role in generating pathogen-induced ROS\textsuperscript{29,32,33}, mediates the ETI-associated ROS. As shown in Fig. 2c, D36E\textit{(avrRpt2)}-induced apoplastic ROS, as detected \textit{in planta} by the H2DCFDA dye, was completely lost in the \textit{rbohd} plant. Consistently, we detected a much compromised ETI resistance against \textit{Pst} DC3000\textit{(avrRpt2)} in the \textit{rbohd} mutant plant (Fig. 2d, Extended Data Fig. 5b). Notably, \textit{rbohd} mutant plants grew to sizes that were similar to wild-type Col-0 plants under optimized growth conditions and, under these conditions, \textit{rbohd} mutant plants showed similar or even slightly enhanced susceptibility to \textit{Pst} DC3000 (Fig. 2d, Extended Data Fig. 5a, b). Altogether, our results suggest a critical dual role of RBOHD in PTI and ETI.

**Requirement of PTI activation of RBOHD for ROS during ETI**

We next assayed the transcript and protein level of RBOHD and found that the transcript (Fig. 2e) and protein level (Fig. 2f) of RBOHD are induced both by D36E and, interestingly, to a much higher level, by D36E\textit{(avrRpt2)} inoculation in Col-0 plant. Surprisingly, the strong induction of \textit{RBOHD} transcript and protein by D36E\textit{(avrRpt2)} occurred at a comparable level in \textit{bbc} mutant plants (Fig. 2e, f). These results indicate that neither \textit{RBOHD} transcript nor RBOHD protein accumulation accounts for the compromised ROS production in the \textit{bbc} mutant and suggests an involvement of PRRs/PTI in post-translational regulation of RBOHD during ETI. Previous studies have reported several classes of kinases, including calcium dependent protein kinases (CPKs) and \textit{Botrytis}-induced kinase 1 (BIK1), involved in phosphorylating RBOHD for ROS production\textsuperscript{32-36}. We examined the ETI-ROS level in the \textit{cpk4/5/6/11} quadruple mutant and \textit{bik1} mutant plants, and found that ETI-associated ROS was reduced in \textit{bik1} mutant but did not seem to be affected in \textit{cpk4/5/6/11} mutant plant (Extended Data Fig. 6), suggesting that BIK1 contributes to the production of ETI-ROS. BIK1 was reported to phosphorylate RBOHD at multiple sites including S39, S343 and S347 during PTI activation\textsuperscript{32,33}. We therefore examined RBOHD phosphorylation levels in protoplasts prepared from Col-0/\textit{DEX::avrRpt2} and \textit{bbc}/\textit{DEX::avrRpt2} plants and transformed with an DNA construct expressing FLAG-RBOHD. PTI and ETI in these protoplasts were activated by \textit{flg22} and DEX, respectively. A 35S promoter was used to express FLAG-RBOHD to ensure similar protein levels during various treatments.
While no phosphorylation at S343/S347 in Col-0 protoplasts was detected 2.5h after treatment with flg22, which is known to induce a PTI-associated transient phosphorylation on RBOHD, DEX alone reproducibly induced a weak phosphorylation of S343/S347 in Col-0 protoplasts 2.5h after treatment (Fig. 2g). Strikingly, a flg22+DEX treatment induced a much stronger phosphorylation on S343/S347 in Col-0 background 2.5h after treatment (Fig. 2g), suggesting a synergistic effect of PTI and ETI in phosphorylating RBOHD. In contrast, no phosphorylation was detected in the bbc background with flg22, DEX or flg22+DEX treatment, confirming the requirement of PRR/co-receptors for RBOHD phosphorylation during ETI. Collectively, our results demonstrate that two classes of immune receptors coordinate to ensure the abundance (i.e. by RPS2) and activity (i.e. by PRRs) of RBOHD for generating robust ETI-ROS. Interestingly, S343/S347 phosphorylation of RBOHD has previously been shown to be important for ETI resistance and restriction of bacterial growth.

PRRs control a subset of ETI-associated transcriptome

The requirement of PRR/co-receptors for activation of RBOHD and a strong up-regulation of RBOHD during ETI (Fig. 2e, f) are intriguing and suggest that ETI may have evolved to co-opt RBOHD and other components of the PTI pathway as part of its mechanism. We therefore examined the expression patterns of other components of the PTI pathway and the rest of Arabidopsis transcriptome by RNAseq (Fig. 3a). Bacteria were infiltrated at a high dose (i.e., ~2x10^7 cfu/mL) and expression was examined at early time points (i.e. 3h and 6h post infiltration) to ensure similar bacterial populations in Col-0 and bbc plants at the sampling times (Extended Data Fig. 7a). We found that, at 3h post infiltration, D36E(avnRpt2) already induced significantly different expression of many genes (i.e., more than 4,000 genes) compared to D36E in Col-0 plant (Extended Data Fig. 7b), suggesting that 3h is sufficient for delivery of AvrRpt2 into the plant cell and triggering strong ETI-associated gene expression. We therefore focused analysis on 3h time point in order to reveal early, and likely more direct, changes of ETI-associated gene expression. Many genes are differentially regulated at this early time point between Col-0 and bbc plants in response to PTI-inducing D36E (Fig. 3b), as expected. Interestingly, these genes show similar expression pattern in Col-0 and bbc plants after D36E(avnRpt2) inoculation to elicit ETI (Fig. 3b), suggesting that ETI can largely restore PTI-
associated global gene expression in the bbc plant. Similar trends were observed for genes associated with salicylic acid, jasmonate and ethylene pathways (Extended Data Fig. 7c-e).

Despite the overall similarity in global gene expression patterns between Col-0 and bbc mutant infected by D36E(\textit{avrRpt2}), we did notice that a subset of 272 genes were differentially expressed (Supplementary table 1). GO analysis of the 209 down-regulated genes (i.e., higher expression in Col-0 compared to that in the bbc mutant) revealed several immune-related terms, such as “response to chitin” and “response to hydrogen peroxide” (Extended Data Fig. 8a). Interestingly, we found that a cluster of \textit{WRKY} genes including \textit{WRKY22/29} and \textit{FRK1} (\textit{Flg22-induced Receptor-like Kinase 1}), which are canonical marker genes of \textit{flg22}-induced PTI pathway\textsuperscript{38}, are down-regulated in the bbc plant during AvrRpt2-triggered ETI (Extended Data Fig. 8b, c). This suggests that, despite the general rescue of global PTI-associated gene expression by ETI in the bbc mutant, the WRKY-FRK1 branch represents a unique immune branch, the activation of which during ETI requires PRR/co-receptors.

**Augmentation of key PTI components by ETI in an SA-independent manner**

We noticed an interesting expression pattern for many PTI signaling genes. As shown in Fig. 3c, while PTI-inducing D36E can induce, to a moderate level, many key PTI components, namely \textit{BAK1}, \textit{BIK1}, \textit{XLG2/AGB1/AGG2}\textsuperscript{39}, \textit{MKK4/5} and \textit{MPK3}, ETI-inducing D36E(\textit{avrRpt2}) induced these genes to a much higher level. Importantly, the strong activation of these PTI components by ETI is independent of PRR/co-receptors, since it occurs in the bbc mutant. Noticeably, BIK1 and some other PBLs, but not PBL1, are strongly induced after D36E(\textit{avrRpt2}) inoculation (Extended Data Fig. 9), suggesting differential contribution of different members of the BIK1/PBL family to ETI. Quantitative RT-PCR was performed to confirm the RNAseq results (Fig. 3d), and western blot further confirmed, at the protein level, the ETI up-regulation of several key components, including \textit{BAK1}, \textit{BIK1} and \textit{MPK3} (Fig. 3e). Our results suggest that part of the AvrRpt2-ETI response is to ensure rapid high-level expression of key components of the PTI pathway, consistent with PTI being an essential component of ETI. We propose that this ETI-mediated up-regulation of PTI components is also likely an important part of a mechanism
to overcome the negative regulation of PTI by endogenous “braking” systems of plants and exogenous pathogen effectors during infections\textsuperscript{2,40}.

A previous study showed that SA treatment could up-regulate PRR protein level\textsuperscript{41}. We therefore tested gene expression in the SA-deficient \textit{sid2} plant and found that PTI components were still up-regulated by D36E(\textit{avrRpt2}) inoculation (Supplementary Fig. 10a). In addition, we examined in our RNAseq dataset the expression pattern of responsive genes to SA and N-hydroxy-pipeolic acid (NHP), which have been shown to function synergistically in plant immunity\textsuperscript{42,43}. Our results showed that both SA- (Extended Data Fig. 7c) and NHP- (Extended Data Fig. 10b) responsive genes had similar transcript level in the Col-0 and \textit{bbc} plants after D36E(\textit{avrRpt2}) inoculation, suggesting intact SA/NHP signaling in the \textit{bbc} mutant during AvrRpt2-ETI. Therefore, ETI seems to “re-enforce” the PTI pathway in a SA/NHP-independent manner.

**Discussion**

Our study reveals a surprising requirement of PRR/co-receptors for effective ETI responses and suggests a model in which ETI co-opts the PTI machinery, including the BIK1-RBOHD module, as an indispensible component (Fig. 4). In particular, we found that PRRs and NLRs, the two primary classes of plant immune receptors, function synergistically to ensure a robust level as well as the fully active status of a key immune component RBOHD, which mediates ETI-ROS generation and full disease resistance. We identified PRR-mediated RBOHD phosphorylation at S343/S347 sites as one of the mechanistic links between PTI and ETI.

Our study sheds light on a long-standing puzzle in the field of plant immunity with respect to the enigmatic similarities in many PTI- and ETI-associated cellular defense features. Our model is supported by a parallel study by Ngou et al. (see back-to-back submission., Manuscript number “BIORXIV/2020/034173”), who focus on the \textit{P. syringae} effector AvrRps4 recognized by TIR-type NLRs (RPS4/RRS1), while we focus on RPS2, a CC-type NLR. Our complementary data suggest conservation of the discovered mechanism for two different types of NLRs, which account for the vast majority of pathogen-sensing NLRs in plants. Intriguingly, synergistic
interaction between surface and intracellular immune receptors in animals and humans has also been reported\(^4^4-^4^7\), suggesting a possible conceptual parallel in immune receptor functions across kingdoms.

The demonstration of PTI as an integral component of ETI has significant implications in understanding how ETI resistance mechanisms prevent pathogen growth. Specifically, several PTI-associated anti-pathogen mechanisms have been described recently, including suppression of bacterial type III secretion\(^4^8-^5^0\), inhibition of bacterial motility by lignification\(^5^1\) and restriction of nutrient acquisition\(^5^2\). Our study suggests that ETI co-opts these PTI mechanisms to halt pathogen growth. Our work likely has significant practical implications as well, as it suggests the possibility for carefully controlled augmentation of PTI components as a new strategy to broadly increase the effectiveness of ETI against numerous plant diseases in crop fields.

References


Methods

Plant materials and growth conditions

Arabidopsis thaliana plants used in this study are in Col-0 ecotype background. The fls2/efr/cerk1 \(^{53}\), bak1/bkk1/cerk1 \(^{16}\), rps2 \(^{54}\), rbohd \(^{29}\), bik1 \(^{55}\), cpk4/5/6/11 \(^{35}\) mutants were reported previously. Plants were grown in potting soil in environmentally-controlled growth chambers, with relative humidity set at 60% and temperature at 22°C with a 12h light/12h dark photoperiod unless stated otherwise. Four- to five-week-old plants were used for all experiments in this study.

To generate the bbc/DEX::avrRpt2 and Col-0/DEX::avrRpt2 transgenic plants, the \(\text{avrRpt2}\) gene was cloned into pBUD-DEX (pBD) vector in the \(Xho\)I/\(Spe\)I restriction enzyme sites, and the expression cassette was introduced into Col-0 or bbc plants by \(\text{Agrobacterium}\)-mediated transformation.

Bacterial disease and HR assays

The \(\text{Pst}\) DC3000 strains carrying \(\text{avrRpt2}\), \(\text{avrRps4}\) and \(\text{avrPphB}\) were published previously \(^{56-58}\). The D36E(\(\text{avrRpt2}\)) strain was generated by transforming the \(\text{avrRpt2}\) expression plasmid into D36E strain by electroporation. For bacterial inoculation, \(\text{Pst}\) strains were cultured in Luria-Marine (LM) medium overnight at 30°C to a cell density of \(\text{OD}_{600}=0.8-1.0\). Bacteria were collected by centrifugation and washed once with sterile water, and adjusted to a cell density of \(\text{OD}_{600}=0.2\). For disease assay, bacterial suspension was further diluted to a cell density of \(\text{OD}_{600}=0.001-0.002\). Bacteria were infiltrated into leaves with a needleless syringe, and inoculated plants were kept under ambient humidity for about 1h to allow evaporation of excess water from the leaf and then covered with a transparent plastic dome to keep high humidity for 3-4 days. For quantification of bacteria, four leaf discs from two different leaves (after surface sterilization) were taken using a cork borer (7.5mm in diameter) as one biological repeat, and 3-4 repeats were taken for each treatment. Leaf discs were ground and diluted in sterile water, and the extraction solutions were then plated on LM agar plates supplemented with rifampicin (at 50mg/L). Colonies were counted with a stereoscope 24h after incubation at 30°C. For HR assay, \(\text{Pst}\) DC3000(\(\text{avrRpt2}\)) suspension was prepared as described above and bacterial suspension at the cell density of \(\text{OD}_{600}=0.2\) was syringe-infiltrated into leaves. Plants were then kept under ambient humidity for about 7h before tissue collapse was recorded.

RIN4 cleavage assays
Arabidopsis plant leaves were infiltrated with *Pst* D36E(*avrRpt2)* (at OD$_{600}$=0.1), and samples were collected at 0, 2, 4, 8h after infiltration by snap-freezing in liquid nitrogen. Three leaves were collected as one biological repeat. Total proteins were extracted in protein extraction buffer (50mM Tris-HCl pH 7.5, 150mM NaCl, 5mM EDTA pH 7.5, 1mM DTT, 1% Triton X-100, 1mM Phenylmethylsulfonyl fluoride) supplemented with 1 x plant protease inhibitor cocktail (Complete EDTA-free, Roche). Cell lysates were centrifuged at 12,000 x g for 15 min at 4°C, and the pellet was discarded. Protein concentration of the supernatant (“total protein extract”) was determined by Bradford protein assay kit (Bio-Rad). An equal amount of total protein was loaded on 12% SDS acrylamide gels (Bio-Rad) for SDS-PAGE. RIN4 protein was detected by anti-RIN4 antibody at a dilution of 1:1000. Total proteins were stained by Coomassie Brilliant Blue (CBB) to show equal loading.

**MAPK kinase activity assay**

Four-week-old plant leaves were infiltrated with *Pst* D36E(*avrRpt2)* (at OD$_{600}$=0.1), and leaves were collected at different time points by snap-freezing in liquid nitrogen. Proteins were extracted in protein extraction buffer (50mM Tris-HCl pH 7.5, 150mM NaCl, 5mM EDTA pH 7.5, 1mM DTT, 1% Triton X-100, 1mM Phenylmethylsulfonyl fluoride) supplemented with 1 x plant protease inhibitor cocktail (Complete EDTA-free, Roche) and 1 x phosphatase inhibitor cocktail (PhosSTOP, Roche). Total protein concentration was determined with Bradford protein assay kit (Bio-Rad). An equal amount of protein was loaded onto 12% SDS-PAGE gel for western blot. Phosphorylated MPK3 and MPK6 proteins were detected by anti-Phospho-p44/42 antibody (Cell Signaling Technology).

**Protein extraction and immunoblotting for PTI signaling components**

Four-week-old plant leaves were infiltrated with sterile water (mock) or different *Pst* strains at OD$_{600}$=0.02, and samples were collected at 0.5, 3, 6, 8h after infiltration. Three to four leaves from different plants were collected as one sample. Protein was extracted using Plasma Membrane Protein Isolation Kit (Invent) according to the manufacturer’s protocol. Concentration of the cytosolic protein was determined with Bradford protein assay kit (Bio-Rad). An equal amount of protein was loaded onto SDS-PAGE gel for western blot. Different PTI components were detected by following antibodies with indicated dilution: anti-RBOHD (Agrisera), 1:1000;
anti-BAK1 (Agrisera), 1:5000; anti-BIK1 (Agrisera), 1:3000; anti-MPK3 (Sigma-Aldrich), 1:2500; anti-MPK6 (Sigma-Aldrich), 1:5000.

**Protoplast transformation and detection of RBOHD phosphorylation**

Protoplasts were prepared from Col-0/DEX::avrRpt2 and bcc/DEX::avrRpt2 plants (4-5 weeks old; grown under 10h light/14h dark photoperiod) and transfected with FLAG-RBOHD plasmid. After overnight incubation to allow protein accumulation, protoplasts were treated with 100nM flg22, 5μM DEX or 100nM flg22+5μM DEX and incubated for 2.5h. Total protein was extracted with protein extraction buffer (50 mM HEPES [pH 7.5], 150 mM KCl, 1 mM EDTA, 0.5% Triton-X100, 1 mM DTT, protease inhibitor cocktail), and then incubated with 50μL anti-FLAG M2 agarose beads (Sigma-Aldrich) for 2 h at 4°C. The bound protein was eluted with 50μL of 0.5mg/mL 3xFLAG peptide for 30 min. RBOHD phosphorylation was detected by immunoblotting with RBOHD-pS343/347 antibody published previously.

**ROS detection Assays**

ROS measurement with luminol-based approach was performed as previously described with minor modification. Briefly, leaf discs of four-week-old *Arabidopsis* plants were harvested using a cork borer (5.5mm in diameter) and floated on 200μL sterile water in a 96-well plate, and then incubated overnight at room temperature under continuous light. On the next day, water was replaced with a solution containing 30mg/L (w/v) luminol (Sigma-Aldrich) and 20mg/L (w/v) peroxidase from horseradish (Sigma-Aldrich) with 100nM flg22 only, 5μM DEX only or 100nM flg22+5μM DEX. The luminescence was detected for 5-6h with a signal integration time of 2min using Varioskan Flash plate reader (Thermo Fisher Scientific). For determining the effects of chemical inhibitors, 10μM diphenyleneiodonium (DPI; Sigma-Aldrich), 15μM salicylhydroxamic acid (SHAM; Sigma-Aldrich) or 1μM sodium azide was added to the elicitation solution at indicated time points and luminescence was recorded as described above.

For detection of ROS production by 2’,7’-Dichlorofluorescein diacetate (H₂DCFDA) under confocal microscopy, plants were infiltrated with *Pst* D36E (OD₆₀₀=0.02) or D36E(avnRpt2) (OD₆₀₀=0.02), air-dried and put back into the plant growth room. ROS was detected at 4-5h post infiltration. Ten μM of H₂DCFDA solution was infiltrated into the leaf and fluorescence signal
was detected 10 min later. Images were captured using a Leica SP8 microscope with a 488 nm
excitation and 501-550 nm emission, and chlorophyll auto-fluorescence was detected at 640-
735 nm.

RNA extraction and qRT-PCR analysis of gene expression

To analyze gene expression levels, four-week-old Arabidopsis plant leaves were infiltrated with
sterile water (mock) or different Pst strains at OD$_{600}$ = 0.04, and then harvested at indicated time
points. Three leaves from different plants were collected as one biological replicate and 4
replicates were collected for each treatment. Samples were frozen and ground in liquid nitrogen.
Total mRNA was extracted using Trizol reagent (Invitrogen) according to the manufacturer’s
protocol. One μg of RNA was used for reverse transcription using the ReverTra Ace® qPCR RT
Master Mix with gDNA remover (TOYOBO). Real-time qPCR analysis was carried out with the
SYBR® Green Realtime PCR Master Mix (TOYOBO) on a CFX real-time machine (Bio-Rad).
Two technical repeats were performed for each sample. The plant U-box gene was used as
reference gene for normalization. Primer sequences for qPCR are listed in Supplementary Table
2.

cDNA library generation and RNAseq

For RNAseq experiments, bacterial inoculation and sample collection were performed as
described above. Two leaves from different plants were harvested as one replicate, and four
biological replicates were collected for each treatment/time point. Total mRNA was extracted
using Trizol reagent (Invitrogen). Total RNA was then treated with DNase I (Invitrogen) to
remove DNA and purified RNA was recovered with RNeasy® MinElute™ Cleanup kit
(QIAGEN) according to the manufacturer’s instructions. Library construction and RNA
sequencing were performed by Novogene company. Briefly, RNA purity and integrity was
examined using the NanoPhotometer® spectrophotometer (IMPLEN) and the RNA Nano 6000
Assay Kit of the Bioanalyzer 2100 system (Agilent Technologies). RNA concentration was
measured with Qubit® RNA Assay Kit in Qubit® 2.0 Flurometer (Life Technologies). One μg
RNA per sample was used as input material for library preparation and sequencing. Sequencing
libraries were generated using NEBNext® Ultra™ RNA Library Prep Kit for Illumina® (NEB),
following the manufacturer’s recommendations and sequenced on Illumina Hiseq platform and
150 bp paired-end reads were generated.
Data analysis of RNA-seq

Clean raw data were obtained by removing reads containing adapter sequences or ploy-N and low-quality reads and were then mapped to the Arabidopsis genome (TAIR10). Gene expression levels were calculated using the TPM method (Transcripts per Kb of exon model per Million mapped reads). Differential expression analysis was performed using the DEGSeq R package (1.18.0). The resulting P-values were adjusted using the Benjamini and Hochberg’s approach for controlling the false discovery rate. Genes with a q-value < 0.05 and log2(Fold change) > 1 found by DESeq were assigned as differentially expressed.

Statistical analysis

All statistical analyses were performed by one-way or two-way analysis of variance (ANOVA) with GraphPad software or two sided student’s t-test with Office Excel software. Each experiment was repeated at least three times and data were represented as the mean ± standard error of mean (s.e.m.) or standard deviation (s.d.) as indicated.

Data availability

All data is available in the main text or the supplementary materials.


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Author contributions: With initial observation of PRR dependency for ETI resistance made by X-F. X. while at Michigan State University, supported by the US National Institute of General Medical Sciences (GM109928), M. Y. and X-F. X. designed all subsequent experiments at Institute of Plant Physiology and Ecology, Chinese Academy of Sciences; M. Y., Z. J., G. B., K. N. and M. L. performed all the experiments described; M. Y. and X-F. X. wrote the paper and S. Y. H. and J-M. Z. edited the paper.

Competing interests: The authors declare no competing interests.

Materials & Correspondence: Correspondence and material requests should be addressed to xinxf@sippe.ac.cn.
## Supplementary Table 2. Primers used in this study

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Fig. 1 | See next page for caption.
PTI-associated PRR/co-receptors are required for ETI responses and resistance. a, Two independent triple PRR/co-receptor mutant plants were hyper-susceptible to Pst DC3000 (avrRpt2) infection. Bacteria were infiltrated into Arabidopsis leaves at OD$_{600}$=0.002 and bacterial populations inside leaves were determined 3 days post infection. b, HR cell death (indicated by leaf collapse in Col-0) was compromised in PRR/co-receptor mutants. Pst DC3000 (avrRpt2) bacteria were infiltrated into Arabidopsis leaves at OD$_{600}$=0.2 and pictures were taken ~7 h after infiltration. Numbers of dead and total infiltrated leaves were counted. c, The triple PRR/co-receptor mutant plants were hyper-susceptible to D36E(avrRpt2) infection. Bacteria were infiltrated into Arabidopsis leaves at OD$_{600}$=0.002-0.004 and bacterial populations inside leaves were determined 3-4 days post infection. Different letters in a, and c, indicate statistically significant differences in bacterial population, as determined by two-way ANOVA (mean ± s.d.; n ≥ 3; p < 0.05). d, e, RIN4 cleavage d and MPK3/6 phosphorylation e in Col-0 and the PRR/co-receptor mutants after D36E(avrRpt2) inoculation. CBB, Coomassie Brilliant Blue staining. An equal amount of total protein was loaded in each lane. f, g, ROS burst detected by luminol-HRP approach in the DEX-inducible avrRpt2 transgenic plant after treatment of 100nM flg22, 5µM DEX, or both. RLUs, relative luminescence units. Total photon counts are calculated from f (n ≥ 27). Different letters indicate statistically significant differences as analyzed by one-way ANOVA (p < 0.05). h, i, ROS burst in Col-0/DEX::avrRpt2 and bbc/DEX::avrRpt2 plants after treatment of 5µM DEX, or 100nM flg22+5µM DEX. Total photon counts are calculated from h (n ≥ 5). Different letters indicate statistically significant differences as analyzed by two-way ANOVA (p < 0.05). Box plots: centre line, median; box limits, lower and upper quartiles; whiskers, highest and lowest data points. Dots indicate individual data points. All experiments were repeated at least three times with similar trends.
Fig. 2 | See next page for caption.
Fig. 2. AvrRpt2-triggered ROS is mediated by RBOHD and requires PRR/co-receptors for activation. a, ROS burst detected with fluorescent dye H$_2$DCFDA in Col-0, bbc, rps2 and rbohd leaves 5h after infiltration of D36E(ivrRpt2) or in Col-0 leaves 5h after infiltration of D36E strain. White arrows indicate the apoplast space in the leaf. Scale bars = 25μm. b, ETI-associated ROS burst is inhibited by DPI, an NADPH oxidase inhibitor. ROS was detected in Col-0/DEX::avrRpt2 plants after treatment of 100nM flg22 and 5μM DEX. Chemical inhibitors (DPI, SHAM or NaN$_3$) were added after the first ROS burst (about 40min after addition of flg22 and DEX). Data are displayed as mean ± s.e.m. (n ≥ 6). c, ROS burst detected with fluorescent dye H$_2$DCFDA in Col-0 and rbohd leaves 5h after infiltration of D36E(ivrRpt2). Scale bars = 25μm. d, The rbohd mutant plant is compromised in ETI resistance against Pst DC3000 (ivrRpt2). Bacteria were infiltrated into Arabidopsis leaves at OD$_{600}$=0.001 and bacterial populations inside leaves were determined 2 days post infection. ***, student’s t-test, two-tailed, p < 0.001. Data are displayed as mean ± s.d. (n = 3). e, RBOHD transcript levels in Col-0 and bbc plants 3h after inoculation of bacterial strains indicated. Data are displayed by mean ± s.e.m. (n ≥ 3). Different letters indicate statistically significant differences, as analyzed by two-way ANOVA (p < 0.05). f, RBOHD protein levels in Col-0 and bbc plants at different time points after inoculation of bacterial strains indicated. The numbers indicate band intensity relative to that of Ponceau S, quantified by ImageJ, and red indicates strong induction. g, Phosphorylation of RBOHD protein at S343/S347 sites requires PRR/co-receptors. FLAG-RBOHD was transformed into protoplasts prepared from Col-0/DEX::avrRpt2 and bbc/DEX::avrRpt2 plants. Protoplasts were treated with elicitors (F, 100nM flg22; D, 5μM DEX; FD, 100nM flg22+5μM DEX) and harvested 2.5h later for FLAG-RBOHD immunoprecipitation. Phosphorylated and total RBOHD proteins were detected by RBOHD pS343/347-specific antibody and FLAG antibody, respectively. All experiments were repeated at least three times with similar trends.
Fig. 3 | ETI upregulates key components of the PTI pathway. 

a, A diagram showing the RNAseq design in this study. 

b, A heat-map of the expression pattern of D36E/PTI-responsive genes in the RNAseq experiment. 

c, A heat map of the expression pattern of PTI pathway genes, showing restoration and up-regulation of expression of major PTI components during AvrRpt2-triggered ETI. 

d, qRT-PCR results of representative PTI pathway genes. Col-0 and bbc plants were infiltrated with different strains indicated, and leaves were harvested 3h post infiltration for transcript analysis (mean ± s.e.m; n ≥ 3). Different letters indicate statistically significant differences as analyzed by two-way ANOVA (p < 0.05). 

e, Protein levels of BAK1, BIK1, MPK3 and MPK6 in Col-0 plants at different time points after inoculation of bacterial strains indicated. BAK1 is detected in the immunoblot of total membrane fraction and other proteins are detected in the immunoblot of total protein extracts. Equal amounts of total proteins were loaded into gel lanes. MPK6 protein is not induced by ETI and serves as an internal control. The numbers indicate band intensity relative to that of Ponceau S, quantified by ImageJ, and red indicates strong induction. All experiments were repeated at least three times with similar trends.
**Fig. 4** A model depicting findings from this study showing PTI as a key component of ETI. Grey color indicates mutated (i.e. FLS2 and BAK1) or inactive (i.e. RBOHD and BIK1) proteins. In wild-type plant, PTI is integrated into ETI in that RPS2 activation leads to protein accumulation of PTI components such as RBOHD and PRR/co-receptors are required for “activating” it by phosphorylation. In the absence of PRR/co-receptors (left panel), although NLR activation strongly induces protein accumulation of PTI components, many of these components, such as BIK1 and RBOHD, are inactive (shown by the grey arrows), leading to compromised ETI resistance.
Extended Data Fig.1 | PRR/co-receptors are required for ETI elicited by different P. syringae avirulent effectors. AvrPphB- and AvrRps4-mediated ETI are also compromised in fec and bbc mutants. Plants were infiltrated with different strains at OD$_{600}$=0.002. Bacterial populations were determined 3 days post inoculation. Different letters indicate statistically significant differences in bacterial populations as determined by two-way ANOVA. (mean ± s.d.; $n \geq 3$; $p < 0.05$). Experiments were repeated at least three times with similar trends.
Extended Data Fig. 2 | The transcript levels of RPS2 are not altered in the fec and bbc mutant plants. RPS2 transcript levels in the fec and bbc mutant plants were similar to those in Col-0 plants after inoculation of bacterial strains indicated. Different letters indicate statistically significant differences as analyzed by two-way ANOVA (mean ± s.e.m; n = 3; p < 0.05). Experiments were repeated at least three times with similar trends.
Extended Data Fig.3 | Characterization of different lines of *bbc/Dex::avrRpt2* plants.

Expression levels of the *avrRpt2* transgene in different transgenic lines 2h after infiltration with 5μM DEX. Different letters indicate statistically significant differences as analyzed by one-way ANOVA (mean ± s.e.m.; *n* ≥ 3; *p* < 0.05). Experiments were repeated at least three times with similar trends.
Extended Data Fig.4| AvrRpt2-triggered ETI-ROS depends on NADPH oxidase. a-c, ROS production in Col-0/DEX::avrRpt2 L1 plants was inhibited by NADPH oxidase inhibitor DPI. Leaf discs were treated with 100nM flg22 and 5μM DEX. DPI, SHAM and NaN₃ were added at the beginning of measurement (mean ± s.e.m.; n ≥ 6). b-c, Total photon counts are calculated from a at the PTI phase (0-30min) or ETI phase (60-200min). Different letters indicate statistically significant difference as analyzed by one-way ANOVA (p < 0.05). Box plots: centre line, median; box limits, lower and upper quartiles; dots, individual data points; whiskers, highest and lowest data points. Experiments were repeated at least three times with similar trends.
**Extended Data Fig.5** The *rbohd* mutant plant is compromised in ETI resistance against *Pst* DC3000(*avrRpt2*). **a**, Appearance of the 5 week-old *rbohd* mutant plants before bacteria inoculation. **b**, Disease symptom of Col-0 and *rbohd* mutant plant 2 days after *Pst* DC3000 and *Pst* DC3000 (*avrRpt2*) infiltration. Experiments were repeated at least three times with similar trends.
Extended Data Fig.6| The AvrRpt2 ETI-associated ROS burst is partially mediated by BIK1. ROS was detected in the bik1 and cpk4/5/6/11 mutant plants by H$_2$DCFDA dye 4.5 h after D36E (avrRpt2) inoculation. Scale bars = 25 μm. Experiments were repeated at least three times with similar trends.
Extended Data Fig. 7 | Transcriptomic analysis of RNAseq experiments. a, Bacterial population in Arabidopsis leaves at 3h or 6h post infiltration. Data are displayed by mean ± s.d. (n = 3). b, A Venn diagram showing numbers of differentially expressed genes (DEGs) 3h after D36E or D36E(avrRpt2) infection in Col-0 plants. c-e, Heat-maps of SA- (c; genes extracted from Karolina et al., 2012\(^{60}\)), jasmonate- (d; genes extracted from Hickman et al., 2017\(^{61}\)) and ethylene-(e; genes extracted from Nemhauser et al., 2006\(^{62}\)) responsive genes.
Extended Data Fig. 8| Genome-wide expression profiling reveals interesting features of gene regulation attributed to PTI and ETI. 

a, GO enrichment analysis of the 209 down-regulated genes in the bbc plant compared to Col-0 plant after D36E(\textit{avrRpt2}) infection (among the total 272 DEGs). b, Heat map of the 272 DEGs in the bbc plant compared to Col-0 plant after D36E (\textit{avrRpt2}) infection, with the canonical PTI pathway genes highlighted in red. c, qRT-PCR of \textit{FRK1} and \textit{WRKY29} expression level in Col-0 and bbc plants 3h after infiltration with different strains or Mock. (mean ± s.e.m.; n = 3; statistical analysis by two-way ANOVA; p < 0.05; different letters indicate statistically significant difference). Experiments were repeated at least three times with similar trends.
Extended Data Fig. 9 | Heat map of BIK1/PBL family gene expression in the RNAseq experiment. Numerical values indicate expression level calculated by TPM (Transcripts per Kb of exon model per Million). Genes labeled in red show significant up-regulation after D36E(AvrRpt2) inoculation, compared to mock and D36E inoculation, in Col-0 and bce plants. Arrows indicate BIK1 and PBL1 genes.
Extended Data Fig. 10 | Up-regulation of key PTI component genes by AvrRpt2-triggered ETI seems to be independent of SA/NHP. a, qRT-PCR analysis of BIK1, XLG2, MKK4, MKK5 and MPK3 expression levels in Col-0 and sid2 plants 3h after infiltration with D36E or D36E(avnRpt2). Different letters indicate statistically significant differences, as analyzed by two-way ANOVA (mean ± s.e.m.; n ≥ 3; p < 0.05). Experiments were repeated at least three times with similar trends. b, Heat-maps of NHP-responsive genes (extracted from Hartmann et al., 201863, defined by genes that are responsive to pipericolic acid and depend on FMO1 for expression) in the Col-0 and bbc plants in our RNAseq experiment.