

Arabidopsis Topless-related 1 mitigates physiological damage and growth penalties of induced immunity

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Summary

- Transcriptional corepressors of the Topless (TPL) family regulate plant hormone and immunity signaling. The lack of a genome-wide profile of their chromatin associations limits understanding of the TPL family roles in transcriptional regulation.
- Chromatin immunoprecipitation with sequencing (ChIP-Seq) was performed on *Arabidopsis thaliana* lines expressing GFP-tagged Topless-related 1 (TPR1-GFP) with and without constitutive immunity via *Enhanced Disease Susceptibility 1 (EDS1)*. RNA-Seq profiling of the TPR1-GFP lines and pathogen-infected *tpl/tpr* mutants, combined with measuring immunity, growth, and physiological parameters was employed to investigate TPL/TPR roles in immunity and defense homeostasis.
- TPR1 was enriched at promoter regions of c. 1400 genes and c. 10% of the detected binding required EDS1 immunity signaling. In a tpr1 tpl tpr4 (t3) mutant, resistance to bacteria was slightly compromised, and defense-related transcriptional reprogramming was weakly reduced or enhanced, respectively, at early (<1 h) and late 24 h stages of bacterial infection. The t3 plants challenged with bacteria or pathogen-associated molecular pattern nlp24 displayed photosystem II dysfunctions. Also, t3 plants were hypersensitive to phytocytokine pep1 at the level of root growth inhibition. Transgenic expression of TPR1 rescued these t3 physiological defects.
- We propose that TPR1 and TPL family proteins function in *Arabidopsis* to reduce detrimental effects associated with activated transcriptional immunity.

Introduction

Plant disease resistance to pathogenic microbes is mediated by cell surface and intracellular immune receptors (Cui et al., 2015; Jones et al., 2016; Albert et al., 2020). Extracellular leucine-rich repeat (LRR) domain receptors recognize pathogen-associated molecular patterns (PAMPs) or host-secreted phytocytokine peptides to confer pattern-triggered immunity (PTI; Albert et al., 2020). Intracellular nucleotide-binding domain/LRR (NLR) immune receptors intercept pathogen virulence factors (called effectors) after their delivery to host cells to produce effector-triggered immunity (ETI). These two receptor systems cooperate to provide robust resistance, often associated with localized host cell death (Ngou et al., 2021; Yuan et al., 2021).

All tested NLR members with N-terminal Toll and interleukin-1 receptor domains (referred to as TIR-NLRs or TNLs) and some cell membrane-resident receptor-like proteins (RLPs) signal via the nucleo-cytoplasmic immunity regulator Enhanced Disease Susceptibility 1 (EDS1; Fradin et al., 2011; Lapin et al., 2020; Dongus & Parker, 2021; Pruitt et al., 2021). EDS1 forms exclusive, functional heterodimers with its sequence-related partners Phytoalexin Deficient 4 (PAD4) and Senescence-associated Gene 101 (SAG101; Wagner et al., 2013; Sun et al., 2021). The EDS1 heterodimers promote timely transcriptional upregulation of defenses in Arabidopsis thaliana (hereafter Arabidopsis), which is necessary for NLR-mediated bacterial resistance (Cui et al., 2018; Mine et al., 2018; Bhandari et al., 2019).

In *Arabidopsis*, WRKY family transcription factors (TFs; Tsuda & Somssich, 2015; Birkenbihl *et al.*, 2017; Zavaliev *et al.*, 2020), Systemic Acquired Resistance Deficient 1 (SARD1) and its homolog Calmodulin-Binding Protein 60-like g (CBP60g; Sun *et al.*, 2015; Ding *et al.*, 2020) have prominent roles in early transcriptional mobilization of defenses. As part of a network with WRKY TFs, CBP60g, and SARD1 help to boost the *Isochorismate Synthase 1 (ICS1)* expression, biosynthesis and signaling of the defense hormone salicylic acid (SA) in response to pathogen attack (Zhang *et al.*, 2010; Zhou *et al.*, 2018). These TFs are further transcriptionally induced in response to salicylic

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acid (SA; Hickman et al., 2019). A myelocytomatosis (MYC) TF, MYC2, controls signaling by the defense hormone jasmonic acid (JA; Lorenzo et al., 2004; Zander et al., 2020). The SA- and JA-dependent signaling branches can antagonize each other, and bacteria employ effector molecules such as coronatine, a structural mimic of JA, to manipulate the hormonal crosstalk (Zheng et al., 2012; Yang et al., 2017). Coronatine-mediated hijacking of JA/MYC2 pathway to dampen SA-dependent defense is blocked in ETI mediated by the TNL pair Resistant to Ralstonia solanacearum 1 (RRS1) and Resistant to Pseudomonas syringae 4 (RPS4) in Arabidopsis (Sohn et al., 2014; Cui et al., 2018; Bhandari et al., 2019). In TNL RRS1-RPS4 ETI, EDS1 enables a timely boost of SA-regulated transcription and suppression of the JA/MYC2-dependent gene expression to restrict bacterial growth (Cui et al., 2018; Bhandari et al., 2019).

Activated defenses can have detrimental effects on plant physiology and growth if they are prolonged or constitutive (Todesco et al., 2010; Ariga et al., 2017; Caarls et al., 2017; van Butselaar & Van den Ackerveken, 2020; Bruessow et al., 2021). DNA methylation and polycomb-dependent H3K27me3 marks, which deplete during plant defense reactions (Dowen et al., 2012; Yu et al., 2013; Dvořák Tomaštíková et al., 2021), small RNAs, and ubiquitin ligase-mediated protein degradation help to limit NLR expression and growth penalties in uninfected plants (Deng et al., 2017; Zervudacki et al., 2018; Copeland & Li, 2019; Huang et al., 2021; Qiao et al., 2021). However, the processes of transcriptional restriction of potentially dangerous induced immunity after pathogen detection are still poorly understood.

Transcriptional corepressors form a layer of gene expression control in eukaryotes. Plant Topless (TPL) and Topless-related (TPR) corepressors resemble Groucho/Tup1 transcriptional corepressors and carry a WD40 repeat C-terminal region and several N-terminal domains (Martin-Arevalillo et al., 2017; Plant et al., 2021). Via the N-terminal domains, TPL/TPRs interact with ethylene response factor (ERF) - amphiphilic repression (EAR) motifs present in multiple TFs (Szemenyei et al., 2008; Causier et al., 2012) and inhibitors of hormone signaling (Pauwels et al., 2010; Ke et al., 2015; Ma et al., 2017; Martin-Arevalillo et al., 2017; Kuhn et al., 2020). Interactions with EAR motifs enable the recruitment of TPL/TPRs into oligomers and complexes with histones, potentially reducing access of TFs to DNA (Ma et al., 2017; Martin-Arevalillo et al., 2017). The N-terminal domain in Arabidopsis TPL further contributes to an oligomerization-independent mode of corepression, likely preventing the engagement of mediator subunits into active transcription complexes (Leydon et al., 2021). TPL/ TPRs also interact with histone deacetylases, providing a mechanism for repression of gene expression by interfering with a transcription-permissive chromatin state (Long et al., 2006; Zhu et al., 2010; Leng et al., 2020). Thus, several molecular mechanisms assist TPL/TPRs corepressor activity.

TPL/TPRs have been implicated in the regulation of plant immunity. First, oomycete and fungal effectors target TPL/TPRs to promote host susceptibility (Harvey et al., 2020; Darino et al., 2021). Second, mutating TPL, TPR1, and TPR4 in Arabidopsis or silencing of TPR1 in Nicotiana benthamiana compromised TNL receptor signaling and an flg22 PAMP-triggered

reactive oxygen species (ROS) burst (Zhu et al., 2010; Zhang et al., 2019; Navarrete et al., 2021). By contrast, Arabidopsis TPR2 and TPR3 were identified as negative regulators of TNL Suppressor of Nonexpressor of Pathogenesis-related 1 (NPR1) Constitutive 1 (SNC1)-conditioned autoimmunity (Garner et al., 2021). Arabidopsis TPR1 is associated with promoters of genes that are downregulated in TNLRRS1-RPS4 ETI (Bartsch et al., 2006; Zhu et al., 2010) and represses the expression of cyclic nucleotide-gated channel (CNGC) genes also known as Defense No Death 1 and 2 (DND1/CNGC2 and DND2/CNGC4; Zhu et al., 2010; Niu et al., 2019). Since these dnd mutants show enhanced bacterial resistance (Clough et al., 2000; Jurkowski et al., 2004), a picture emerged in which TPR1 promotes TNL ETI by limiting the expression of negative regulators of defense (Zhu et al., 2010). However, the lack of a genome-wide profile of TPL/TPR chromatin associations leaves other functions of these corepressors in defense signaling unclear.

Here, using chromatin immunoprecipitation with sequencing (ChIP-Seq), we examined genome-wide TPR1-chromatin associations conditional on *EDS1*-controlled immunity in *Arabidopsis* lines expressing *pTPR1:TPR1-GFP*. We further examined RNA expression profiles and physiological phenotypes of wild-type and *tpr1 tpl tpr4 (t3)* mutant plants during bacterial infection. Taken together, our data suggest that TPL/TPR transcriptional corepressors operate at the chromatin not only to assist in the restriction of pathogen growth but also to mitigate deleterious effects of induced immunity on plant health.

Materials and Methods

Plant materials and growth conditions

Arabidopsis thaliana (L.) Heynh. accession Col-0 tpr1 single mutant, tpr1 tpl tpr4 (t3) triple mutant, pTPR1:TPR1-GFP Col-0 (TPR1 Col), and pTPR1:TPR1-HA Col-0 stable transgenic lines were described previously (Zhu et al., 2010). pTPR1:TPR1-GFP eds1-2 (TPR1 eds1) and pTPR1:TPR1-GFP sid2-1 (TPR1 sid2) lines were generated by crossing TPR1 Col (Zhu et al., 2010) with Col-0 eds1-2 (Bartsch et al., 2006) and Col-0 sid2-1 (Wildermuth et al., 2001), respectively. Complementation tpr1 tpl tpr4 pTPR1:TPR1-GFP lines were generated by floral dipping of t3 with Agrobacteria GV3101 pMP90 pSoup carrying pCAMBIA1305-TPR1-GFP (Zhu et al., 2010). The coi1-41 mutant is described in Cui et al. (2018). The mutant eds1-2 (Bartsch et al., 2006) was used throughout the study, with the exception of root growth inhibition and MAPK assays, where the eds1-12 line (Ordon et al., 2017) was used. The mutant eds1-12 pad4-1 sag101-3 (eps) is described in Ordon et al. (2017) and Pruitt et al. (2021). The fls2 (SAIL_691C4), rlp23-1, and pepr1-1 pepr2-3 mutants are described earlier (Zipfel et al., 2004; Krol et al., 2010; Albert et al., 2015). Oligonucleotides for genotyping are shown in Supporting Information Table S1. For bacterial infection assays, plants were grown under a 10 h light period (c. 100 μ mol m⁻² s⁻¹) and 22°C: 20°C, day: night temperature regime with 60% relative humidity. For transformation and selection of combinatorial mutants, plants were grown under

22 h light (c. 100 μ mol m⁻² s⁻¹) and a 22°C: 20°C, day: night temperature regime with 60% relative humidity.

Immunoblot analyses

For immunoblotting of TPR1-GFP, total protein extracts were prepared by incubating liquid nitrogen-ground samples (c. 50 mg) in 2× Laemmli loading buffer for 10 min at 95°C. Samples were centrifuged for 1 min at 10 000 g to remove cell debris before gel loading. Proteins were separated by 10% (v/w) SDS-PAGE (1610156; Bio-Rad, Hercules, CA, USA) and transferred to a nitrocellulose membrane (0600001; GE Healthcare Life Sciences, Chicago, IL, USA). α-GFP antibodies (no. 2956; Cell Signaling Technology, Danvers, MA, USA, or no. 11814460001; Roche, Basel, Switzerland) in combination with HRP-conjugated anti-rabbit or antimouse secondary antibodies (A9044 or A6154; Merck, Darmstadt, Germany) were used. In MAPK3/6 phosphorylation assays, seedlings were treated for 15 and 180 min with pep1 or milliQ water (mQ, mock) as a negative control. Proteins were extracted with a buffer containing 50 mM Tris pH 7.5, 200 mM NaCl, 1 mM EDTA, 10 mM NaF, 2 mM sodium orthovanadate, 1 mM sodium molybdate, 10% (v/v) glycerol, 1 mM AEBSF, 0.1% Tween-20, 1 mM dithiothreitol, 1× protease inhibitor cocktail (11836170001; Roche), and 1× phosphatase inhibitor cocktail (4906845001; Roche). Extracts were resolved on 8% (v/w) SDS-PAGE (1610156; Bio-Rad) and transferred onto a nitrocellulose membrane (0600001; GE Healthcare Life Sciences). Primary antibody against phospho-p44/42 MAP kinase (#9101; Cell Signaling Technologies) was used in combination with HRP-conjugated anti-rabbit as a secondary antibody (A6154; Merck). Signal detection was performed using Clarity and Clarity Max luminescence assays (1705061 and 1705062; Bio-Rad). For loading control, membranes were stained with Ponceau S (09276-6X1EA-F; Merck).

Salicylic acid quantification

Quantification of free SA was done as described (Straus et al., 2010) with a chloroform/methanol/water extraction containing SA-d₄ (CS04-482_248; Campro Scientific, Berlin, Germany) as internal standard. After phase extraction, drying of polar phase, dissolving in sodium acetate (pH 5.0), uptake in ethyl acetate: hexane (3:1), and derivatization, 1 µl sample was injected into a gas chromatograph coupled to a mass spectrometer (GC–MS; Agilent, Santa Clara, CA, USA) on an HP-5MS column (Agilent). Masses of SA-d₄ (m/z 271) and SA (m/z 267) were detected by selected ion monitoring and quantified using the CHEMSTATION software (Agilent).

Chla fluorescence and chlorophyll quantification

Maximum quantum efficiency (F_v/F_m) of photosystem II (PSII) and the effective efficiency (ϕ PSII) in Col, tpr1, t3, and eds1 leaves were determined after syringe infiltration of Pst (OD₆₀₀ = 0.005) by Chla fluorescence analysis using a MINI-PAM fluorimeter (Walz, Effeltrich, Germany). Measurements of three to four leaves from independent plants were performed at each timepoint in a

randomized and rotating order between 13:00 and 15:00 h on Days 0-4 after inoculation (10:00-11:00 h). Mock (10 mM MgCl₂)infiltrated leaves from different plants were measured as controls. To determine the maximum quantum yield $(F_v/F_m = (F_m - F_0)/$ $F_{\rm m}$; Baker, 2008), plants were first dark-acclimated for 20 min. The operating PSII efficiency of photosystem II (φ PSII = $(F_m' - F)/F_m'$; Baker, 2008) was determined with 12 saturating light flashes (c. 1300 μ mol photons m⁻² s⁻¹) at intervals of 20 s under actinic light intensity of c. 216 µmol photons m⁻² s⁻¹. Data from three independent experiments were combined, statistically analyzed using ANOVA and Tukey's HSD test ($\alpha = 0.05$), and plotted using the 'ggline' function in the 'GGPUBR' R package. In Chla fluorescence assays with nlp24 treatment, nlp24 (AIMYAWYFPKDSPM LLMGHRHDWE, crude peptide; Genscript, Piscataway, NJ, USA; dissolved in DMSO) was applied by leaf infiltration at a final concentration of 5 µM (10 mM MgCl₂, 0.05% DMSO). Data from four independent experiments were combined and statistically analyzed using ANOVA and Tukey's HSD test $(\alpha = 0.05)$. Total leaf chlorophyll (a + b) contents were determined at 3 d after syringe infiltration with P. syringae pv tomato DC3000 bacteria (OD $_{600} = 0.005$) or mock (10 mM MgCl₂) treatment. The chlorophyll content in each sample was calculated as a mean of three leaf disks (diameter 8 mm) and analyzed according to Porra et al. (1989). Three independent experiments were performed and pooled for the statistical analysis keeping experiment as a factor in the ANOVA model (Tukey's HSD $\alpha = 0.05$; n = 15).

Root growth inhibition assay

Root growth inhibition assays with pep1 and flg22 were performed as described (Igarashi et al., 2012), with adjustments. Seeds were surface-sterilized and transferred into 48-well plates (one seed per well). Each well was supplied with 200 μl of 0.5× MS (including vitamins and MES, pH 5.4; M0255; Duchefa Biochemie, Haarlem, the Netherlands) and 5% (w/v) sucrose. flg22 (RP19986; GenScript) and Atpep1 (referred to as pep1 in this study; ATKV-KAKQRGKEKVSSGRPGQHN; GenScript) peptides were dissolved and administered as solutions in mQ water (final concentrations: flg22 - 100 nM, pep1 - 50 or 200 nM). Sterile mQ was added as a mock control. Root length was measured at 10 d using IMAGEJ software. Root growth inhibition (RGI) index was quantified as a ratio of the root length in flg22 or pep1 treatment to the mean of mock-treated plants. Data from independent experiments were combined and statistically analyzed using ANOVA (experiment as a factor) and Tukey's HSD test.

Details on the TPR1-GFP ChIP- and RNA-Seq procedures and data analysis, as well as other assays, are in Methods \$1.

Results

Arabidopsis TPR1 Col displays constitutive transcriptional immunity

To investigate the role of TPR1 in plant immunity, we used an *Arabidopsis* Col-0 line expressing *TPR1-GFP* under control of its

2 kb upstream sequence (pTPR1:TPR1-GFP; hereafter TPR1 Col) and displaying EDS1- and TNL SNC1-dependent constitutive immunity and SA accumulation (Zhu et al., 2010). We introduced a null eds1 (eds1-2) or ics1 (sid2-1) mutation into TPR1 Col to test TPR1-GFP functions without EDS1- or ICS1/ SA-dependent signaling (Wildermuth et al., 2001; Bartsch et al., 2006). While TPR1-GFP accumulation was similar in all three lines (Fig. 1a), stunting of 5-6-wk-old TPR1 Col plants was reduced in TPR1 eds1 but not in TPR1 sid2 (Zhu et al., 2010; Figs 1b, S1a). Enhanced resistance of TPR1 Col to virulent P. syringae pv tomato DC3000 (Pst) bacteria (Zhu et al., 2010) was abolished in TPR1 eds1 and partially compromised in TPR1 sid2 plants (Zhu et al., 2010; Fig. S1b). TPR1 eds1 and TPR1 sid2 plants had c. sevenfold lower SA levels than TPR1 Col (Fig. S1c). These results suggest that constitutive defense in TPR1 Col is mediated primarily by an SA-independent branch of EDS1 signaling, consistent with TPR1 Col autoimmunity involving TNL SNC1 (Zhu et al., 2010), which promotes SAindependent signaling (Zhang et al., 2003; Zhu et al., 2010).

RNA-Seq analysis of 5-6-wk-old TPR1 Col, TPR1 sid2, TPR1 eds1, and wild-type Col plants showed that EDS1 signaling impacted 61% of genes that are differentially expressed between TPR1 Col and Col (Table S2; 942/1549, $|\log_2 FC| \ge 2$, FDR \leq 0.05; Fig. S1d). By contrast, the *sid2* mutation affected the expression of c. 10% differentially expressed genes (DEGs; Table S2; 153/1549, $|log_2FC| \ge 2$, $FDR \le 0.05$). The 2194 DEGs between Col, TPR1 Col, TPR1 sid2, and TPR1 eds1 fell into 13 groups in hierarchical clustering of log₂-transformed gene expression changes relative to Col (Fig. 1c; Table S3). Clusters 8 and 9 were skipped because their small size (< 20 genes) could affect the validity of statistical tests, but they are shown in Table S3. Cluster #1, with 524 genes induced in a TPR1/EDS1dependent manner, was strongly enriched for Gene Ontology (GO) terms linked to EDS1- and SA-dependent immune responses (Fig. 1c; Table S4). By contrast, cluster #10, with 394 genes suppressed in TPR1 Col (Fig. 1c), was enriched for genes linked to microtubule-based dynamics and cell cycle regulation (Table S4). These data show that TPR1-GFP constitutive immunity involves EDS1-dependent transcriptional reprogramming.

We tested whether the TPR1 Col transcriptome aligns with gene expression profiles of PTI and ETI. For this, we crossreferenced DEGs from the 13 clusters above (Fig. 1c; Tables S2, S3) with RNA-Seq datasets for (1) Col inoculated with Pst avrRps4 triggering an ETI^{RRS1-RPS4} (Bhandari et al., 2019) and (2) Col treated with the bacterial PAMP flg22 (Birkenbihl et al., 2017; Figs 1d, S1e). Genes in clusters 1, 3, 4, 6, and 7 that were upregulated in TPR1 Col vs Col (Fig. 1c) were also induced by Pst avrRps4 or flg22 treatments (Fig. 1d). Similarly, repressed clusters in TPR1 Col (#10, #11, Fig. 1c) were downregulated by these treatments (Fig. 1d). Accordingly, there was a significant overlap of DEGs induced in TPR1 Col and treatments with flg22 or Pst avrRps4 (Fig. 1e, Fisher's exact test P < 0.05). We concluded that the TPR1 Col line displays constitutive transcriptional immunity and that TPR1 Col and TPR1 eds1 are suitable backgrounds to measure immunity-dependent and immunity-independent TPR1-chromatin associations.

TPR1 binds to promoters of genes upregulated in immunity activated tissues

We performed a ChIP-Seq analysis on leaves of 5-6-wk-old TPR1 Col and TPR1 eds1 plants (Fig. 2; Methods S1) using an input control for peak calling. A line expressing pTPR1:TPR1-HA in Col, which has a similar constitutive immunity phenotype as TPR1 Col (Zhu et al., 2010), was included as an additional control for peak calling. In TPR1 Col, 1531 TPR1-GFP chromatin binding sites corresponded to 1441 genes (Table \$5). Most peaks (723/1531, 47%) mapped to 1-kb gene sequences as indicated by a metaplot analysis (Table S5; Fig. 2a,b). The results are consistent with TPR1 acting as a transcriptional corepressor at the promoters of CNGC2 and CNGC4 (Niu et al., 2019). TPR1-bound genes showed enrichment of GO terms linked to defense and SA signaling as well as developmental processes (Table S6, FDR \leq 0.05; Figs S2-S4), as expected from the TPR1 Col enhanced defense and perturbed growth phenotypes (Fig. 1b,c).

In TPR1 eds1, which lacks constitutive immunity (Figs 1b,c, S1), we detected 614 TPR1-GFP-binding sites corresponding to 623 genes (Table \$7; Fig. 2c). While the reduced number of peaks in TPR1 eds1 did not affect TPR1 distribution across genomic fractions relative to TPR1 Col (Fig. 2b,c; Table S7), the proportion of defense-related GO terms enriched among TPR1-GFP-bound genes plummeted in TPR1 eds1 relative to TPR1 Col (Table S8). Hence, TPR1-chromatin association with defense-related genes is likely enhanced in immune-activated shoot tissues. To assess this further, we compared TPR1chromatin associations in TPR1 Col and TPR1 eds1 using a peak calling-independent method implemented in diffReps (Shen et al., 2013). This analysis showed that TPR1-GFP enrichment was stronger in TPR1 Col relative to TPR1 eds1 at 247 genes (Gtest, 1.5 times difference, FDR ≤ 0.05; Table \$9), supporting stronger TPR1 binding at these loci in immune-activated TPR1 Col. No ChIP peaks were called for 150 (61%) of these genes in TPR1 eds1 (Tables S5, S7). Notably, 66 of the 247 differentially TPR1-bound genes (27%), including ICS1, cysteine-rich receptor-like kinases, and WRKYTFs (Fig. S2), were more highly expressed in TPR1 Col compared with TPR1 eds1 (Table S2, $\log_2 FC \ge 1$, FDR ≤ 0.05 ; Fig. 2d). Only 10 genes from the above set of 247 (c. 4%) were downregulated in TPR1 Col compared with TPR1 eds1 (Table S2, $\log_2 FC \le 1$, FDR ≤ 0.05 ; Fig. 2d). In summary, TPR1 binds to c. 1400 genes mainly at promoter regions and c. 11% of detected TPR1 binding (150/1441 genes) is conditional on *EDS1*-dependent immunity.

We further tested whether *EDSI*-dependent TPR1-chromatin associations correlate with transcriptional reprogramming during defense. The set of 247 genes with a stronger TPR1-GFP signal in *TPR1* Col vs *TPR1 eds1* (Tables S9, S10) was induced after treatments with PAMP flg22 or *Pst avrRps4* (Fig. 2e, boxplots with orange shadowing). Conversely, the expression of 74 genes with lower TPR1-GFP enrichment in *TPR1* Col vs *TPR1 eds1* (Tables S9, S10) was unaltered or reduced in these treatments (Fig. 2e, boxplots with green shadowing). Similarly, genes bound by TPR1-GFP in *TPR1* Col were specifically enriched for genes

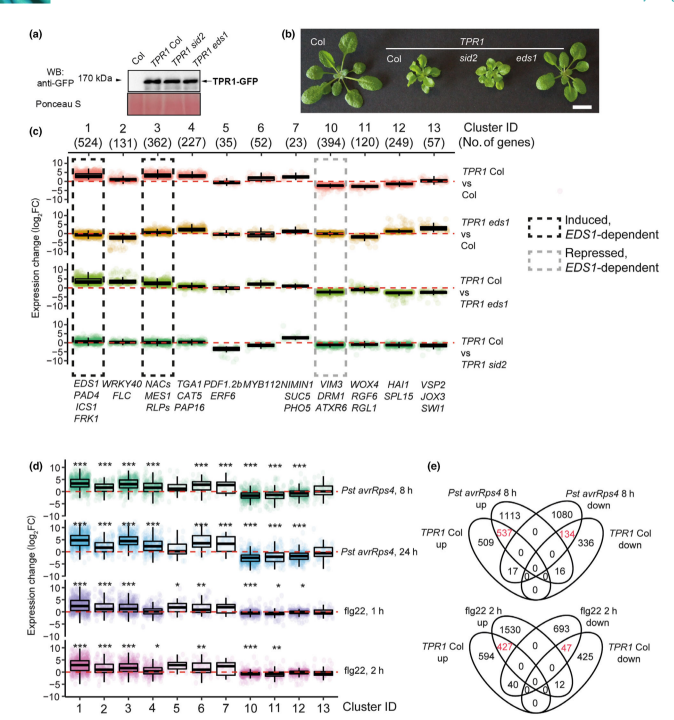


Fig. 1 Defense-related *EDS1*-dependent transcriptional reprogramming in *TPR1* Col line. (a) TPR1-GFP steady-state accumulation in 5–6-wk-old *Arabidopsis* Col-0 (Col), *sid2*, and *eds1* mutant plants expressing *pTPR1:TPR1-GFP* (*TPR1* col, *TPR1* sid2, and *TPR1* eds1). The transgenic lines show similar levels of TPR1-GFP protein. Col was used as a negative control. Ponceau S staining indicates similar loading. The experiment was repeated three times with similar results. (b) Dwarfism in *TPR1* Col depends on functional *EDS1*. Col is shown on the left for comparison. Bar, 1 cm. (c) Log₂-transformed relative expression values of clusters of genes differentially expressed in Col, *TPR1* eds1, and *TPR1* sid2 relative to *TPR1* Col or Col. Each dot corresponds to a single gene. Size of the cluster is given in parentheses. Genes in clusters 1 and 3 are upregulated (box with black dashed line) while cluster 10 (box with gray dashed line) is downregulated in an *EDS1*-dependent manner in *TPR1* Col relative to Col. Clusters 8 and 9 were skipped because their small size (< 20 genes) could affect validity of statistical tests, but they are shown in Supporting Information Table S3. Names of selected genes from the clusters are in italics. (d) Expression changes for genes in clusters from (c) in Col plants treated with *Pseudomonas syringae* pv *tomato* DC3000 *avrRps4* (*Pst avrRps4*) or flg22 at the indicated time points (Birkenbihl *et al.*, 2017; Bhandari *et al.*, 2019). The values are log₂-transformed fold changes (FC) relative to mock or untreated Col plants.

Asterisks indicate that the mean log₂FC value in the cluster differs from 0 based on two-sided *t*-test followed by Bonferroni correction for multiple testing.

*, P < 0.05; ***, P < 0.01; ****, P < 0.001. (e) Venn diagrams showing the overlap between DEGs in *TPR1* Col (vs Col) and DEGs after flg22 and *Pst avrRps4*. Numbers are in red when Fisher's exact test shows statistically significant overlap (*P* < 0.05, Bonferroni correction).

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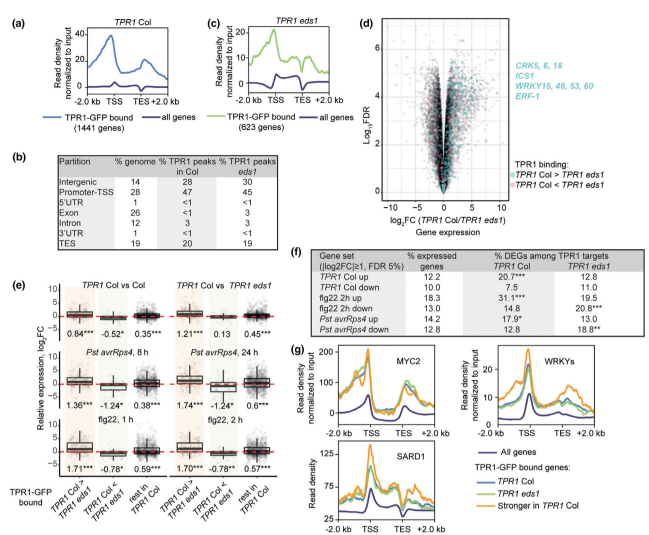


Fig. 2 Arabidopsis TPR1-chromatin association partially depends on EDS1-controlled immune signaling. (a-c) Metaplots of ChIP-Seq TPR1-GFP enrichment profiles at the chromatin in TPR1 Col (a) and TPR1 eds1 (c) and distribution of TPR1 peaks over genome partitions (b). TPR1-GFP binds 1441 genes in TPR1 Col and 623 genes in TPR1 eds1. The ChIP-Seq read density for TPR1-GFP was normalized to input via subtraction. The dark blue lines represent TPR1-GFP chromatin-binding profiles averaged across all annotated genes in Arabidopsis (TAIR10). TES, transcription end site; TSS, transcription start site. (d) Volcano plot displaying the relationship between EDS1-dependent TPR1-chromatin associations and the EDS1-dependent gene expression regulation in TPR1 Col. Significance of difference in the TPR1-GFP enrichment in TPR1 Col and TPR1 eds1 was assessed with diffReps (≥ 1.5 times, G-test, FDR ≤ 0.05). Genes with stronger enrichment of TPR1-GFP in TPR1 Col. than in TPR1 eds1 (blue dots) tend to have higher gene expression in TPR1 Col. Selection of these genes is shown in blue text. (e) Log₂-scaled fold change of the relative expression of TPR1-GFP-bound genes in TPR1 Col vs Col and TPR1 Col vs TPR1 eds1, for treatment with Pseudomonas syringae pv tomato DC3000 (Pst) avrRps4 (8 and 24 hpi vs 0 hpi) or flg22 (1 and 2 hpi vs 0 hpi) (Birkenbihl et al., 2017; Bhandari et al., 2019). Boxplots for genes showing stronger TPR1-GFP enrichment in TPR1 Col vs TPR1 eds1 (shown as 'TPR1 Col > TPR1 eds1', n = 247) are shaded in orange. Green shadowing marks a set of genes with weaker TPR1-GFP signal in TPR1 Col vs TPR1 eds1 ('TPR1 Col < TPR1 eds1', n = 74). Categorization is based on the diffRep results ($log_2|G| > 0.58$ or 1.5 times difference at $P_-adj \le 0.05$. Genes that do not fall in these two categories are denoted as 'rest in TPR1 Col' (n = 1253). The total number of genes in (e) 1574 differs from 1441 in (a) because diffReps works independently of peak calling while annotations in (a) are based on the peak calling. Genes with higher TPR1-GFP enrichment in TPR1 Col show transcriptional upregulation after flg22 and Pst avrRps4 treatments. Numbers below boxplots indicate mean log₂FC; its difference from 0 was assessed with twosided t-test; Bonferroni-corrected P-values: *, P < 0.05; ***, P < 0.01; ***, P < 0.001. Elements of boxplots: first quartile – minima, third quartile – maxima, median – central line; whiskers extend to the minimum and maximum values but not further than 1.5 interquartile range from the respective minima or maxima of the boxplot. Each dot represents a gene. (f) TPR1-GFP-associated loci in TPR1 Col are enriched for genes upregulated in TPR1 Col, after treatments with flg22 (2 h) and $Pst\ avrRps4$ (8 h) (same as in e). Fisher's exact test P-value after Bonferroni correction: *, P < 0.05; **, P < 0.01; ***, P < 0.001. Number of expressed genes depends on the threshold applied during the differential gene expression analysis to remove low-expressed genes (TPR1 Col – 18 884 genes; flg22 - 21 098; Pst avrRps4 - 20 573). (g) Distribution of ChIP-Seq signal for MYC2 (Wang et al., 2019), WRKY (Birkenbihl et al., 2018), and SARD1 (Sun et al., 2015) transcription factors (TFs) across genes bound by TPR1-GFP in TPR1 Col (light blue), TPR1 eds1 (green) and genes bound stronger by TPR1-GFP in TPR1 Col than in TPR1 eds1 (orange). TF-chromatin binding profiles averaged across all annotated genes in Arabidopsis genome (dark blue) serve as a baseline. MYC2, WRKY TFs, and SARD1 are strongly enriched in promoters of genes bound by TPR1-GFP in TPR1 Col and TPR1 eds1. ChIP-Seq data for SARD1 (Sun et al., 2015) did not have input samples and therefore were not normalized, but specificity of SARD1 binding was demonstrated in the original study. ChIP-Seq for MYC2 (Wang et al., 2019) and WRKY TFs (Birkenbihl et al., 2018) were normalized to the input via subtraction.

upregulated in *TPR1* Col (compared with Col) and in response to flg22 or *Pst avrRps4* (Fig. 2f; Table S11). Together, these observations suggest *EDS1*-dependent TPR1 binding to a set of genes that are upregulated in PTI and ETI.

TPR1 ChIP-Seq identifies known TPR1 and TPL targets

We detected TPR1-GFP enrichment at nine of 12 downregulated genes in TNL^{RRS1-RPS4} ETI (Fig. S3) that were identified as TPR1-bound targets in a previous ChIP-qPCR study using the TPR1-HA Col transgenic line (Zhu et al., 2010). Genes with TPR1-GFP enrichment include DND1 and DND2 (Fig. S3) encoding CNGC2 and 4, respectively, which are required for calcium-dependent immunity responses in PTI and ETI (Clough et al., 2000; Jurkowski et al., 2004; Tian et al., 2019). TPR1-GFP binding at these loci was not obviously altered in TPR1 eds1 (Fig. S3), indicating immunity status-independent association of TPR1 with promoters of these nine genes. Since TPL/TPRs have redundant functions (Zhu et al., 2010; Harvey et al., 2020; Plant et al., 2021), we expected overlap in binding targets between TPL and TPR1. Indeed, TPR1-GFP was enriched at several TPL targets found with ChIP-qPCR such as Constans (Goralogia et al., 2017), Apetala 3 (Gorham et al., 2018), Circadian clock associated 1, Leafy, and others (Lee et al., 2020) in both TPR1 Col and TPR1 eds1 (Fig. S4). Hence, our TPR1-GFP ChIP-Seq profiles provide a genome-wide resource for identifying TPL/ TPR chromatin targets.

TPR1 shares binding targets with MYC2, SARD1 and WRKY TFs

The genome-wide profiles of TPR1-chromatin associations in immune-activated and nonactivated leaf tissues prompted us to investigate whether certain DNA motifs correlate with TPR1 binding. A de novo motif search revealed strong enrichment of the GAGA motif (C-box) under TPR1 peaks in TPR1 Col and TPR1 eds1 (Fig. S5a). The G-box (CACGTG) bound by MYC2 and other bHLH TFs was also over-represented under TPR1-GFP peaks in TPR1 eds1 (Fig. S5a). We validated this signature by reanalyzing published MYC2 ChIP-Seq profiles (Figs 2g, S5b). A MYC2 ChIP signal (Wang et al., 2019) was similarly enriched at promoters of genes bound by TPR1-GFP in TPR1 Col and TPR1 eds1 (Fig. 2g). Also, TPR1-bound genes showed statistically significant enrichment of reported MYC2 targets (Van Moerkercke et al., 2019; Zander et al., 2020; Fig. S5b). Our de novo motif searches did not find evidence for the enrichment of W-box 'TTGACY' bound by WRKYs (Ciolkowski et al., 2008) or the 'GAAATTT' element bound by SARD1 (Sun et al., 2015). Considering the importance of these TFs in immune responses, we specifically examined the distribution of WRKY and SARD1 TFs binding at TPR1-GFP-bound genes using available ChIP-Seq data (Sun et al., 2015; Birkenbihl et al., 2018; Figs 2g, S5c,d). The metaplot and enrichment analyses for sets of genes associated with TPR1-GFP and TF peaks revealed that WRKY TFs and SARD1 binding sites strongly overlap with those for TPR1-GFP (Figs 2g, S5c,d). These results

suggest that TPR1 shares some *in vivo* binding targets with MYC2, SARD1, and WRKY TFs.

TPL/TPRs suppress prolonged expression of TNL^{RRS1-RPS4} ETI-induced genes

To explore the functions of TPR1 and other TPL/TPRs in pathogen defense, we infiltrated Arabidopsis tpr1 single and tpr1 tpl tpr4 triple (t3) mutants with avirulent (TNL RRS1-RPS4inducing) Pst avrRps4 or virulent Pst (EV) bacteria alongside Col and hyper-susceptible Col eds1-2 (eds1). As expected for ETI, Col showed stronger restriction of Pst avrRps4 than Pst growth $(0.7 \log_{10}; \text{Tukey's HSD } P = 0.00007) \text{ (Fig. 3a,b)}$. The difference in colony-forming units (CFU) counts is similar to earlier studies from our laboratory (Lapin et al., 2019; Sun et al., 2021) but lower than in other studies (Saucet et al., 2015; Ngou et al., 2021), suggesting that experimental conditions (e.g. humidity) influence absolute differences in CFU of these strains in Col. Growth of Pst avrRps4 and Pst in tpr1 was not different from Col, and t3 had two- to threefold increased CFU counts at 3 d (0.3–0.4 on \log_{10} scale, P = 0.277 and 0.026 in assays with Pst avrRps4 and Pst, respectively; Fig. 3a,b).

Arabidopsis TPL represses MYC2 (Pauwels et al., 2010) which, when activated by bacterial coronatine, antagonizes EDS1- and ICSI/SA-dependent bacterial resistance (Zheng et al., 2012; Cui et al., 2018; Bhandari et al., 2019). We therefore tested whether defects of tpr1 and t3 mutants in bacterial resistance are masked by coronatine. For that, tpr1 and t3 plants were infiltrated with coronatine-deficient Pst Δcor avrRps4 or Pst Δcor (Fig. 3c,d). A mutant of the coronatine insensitive 1 (COI1) JA coreceptor was included as a negative control (Zheng et al., 2012). In line with earlier studies, Col displayed bacterial coronatine-dependent susceptibility compared with coil (Fig. 3b,d), and Pst coronatinepromoted virulence was no longer detected in avrRps4-activated TNLRRS1-RPS4 ETI (Cui et al., 2018; Bhandari et al., 2019; Fig. 3a,b,d). These results confirm that TNLRRS1-RPS4 ETI was functional in our assays. In the absence of coronatine, differences in bacterial growth between Col and the t3 mutant increased to 0.5–0.6 on log_{10} scale (P=0.012 and 0.0002 for Pst Δcor avrRps4 or Pst $\triangle cor$, respectively; Fig. 3c,d). We concluded that TPR1/TPL contribute to Arabidopsis basal resistance and ETI against Pst (Zhu et al., 2010), but this is largely masked by bacterial coronatine virulence.

We performed RNA-Seq on leaves of the tpr1 and t3 mutants alongside Col and eds1 using Pst avrRps4 since the timing of major EDS1-dependent transcriptional reprogramming between 4 and 24 hpi is well established for this strain (Bhandari et al., 2019; Saile et al., 2020; Sun et al., 2021). Leaves of 5–6-wk-old plants were infiltrated with Pst avrRps4 and samples collected at 0 (c.5 min), 8, and 24 hpi (Table S12). As expected, the number of transcriptionally induced genes was higher in Col compared with eds1 at 8 (2097 genes) and 24 (1289 genes) hpi (Table S12, $log_2FC \ge 1$, $FDR \le 0.05$). Surprisingly, only one DEG was detected between Col and tpr1 or t3 mutants at these time points (Table S12, $log_2FC \ge 1$, $FDR \le 0.05$). This was a gene of

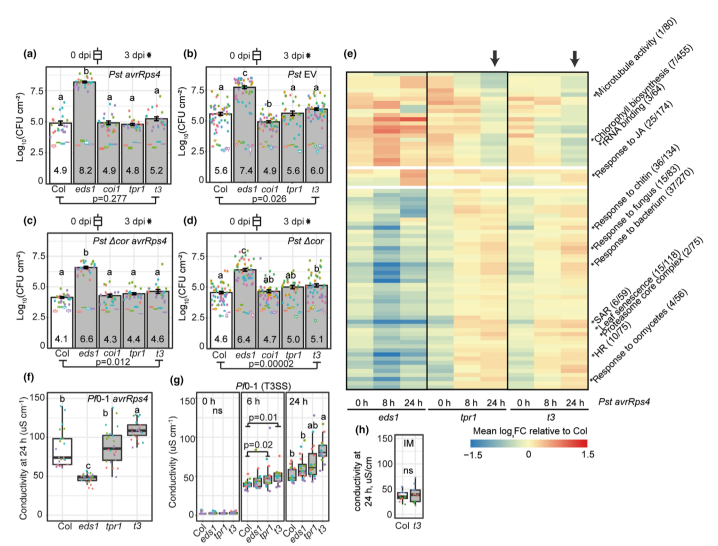


Fig. 3 Arabidopsis TPL/TPRs have a dual role during bacterial resistance responses. (a-d) Titers of Pseudomonas syringae pv tomato DC3000 (Pst) avrRps4 (a), Pst (b), Pst avrRps4 \(\Delta\)cor (c), Pst \(\Delta\)cor (d) bacteria in indicated \(Arabido\)psis mutants relative to Col plants (3 d after syringe infiltration, 3 d postinoculation (dpi), OD₆₀₀ = 0.001). eds1 mutant served as a susceptibility control, and the coi1 mutant – as a readout for the coronatine-promoted susceptibility. The tpr1 tpl tpr4 (t3) mutant showed higher, but not significantly increased levels of the Pst avrRps4 and Pst growth compared with Col (Tukey's HSD, $\alpha = 0.001$; n = 22 from four independent experiments with *Pst avrRps4* and n = 46 from eight independent experiments with *Pst*). *Pst* grows to higher densities than Pst avrRps4, Pst Δcor , or Pst Δcor avrRps4 in Col plants (codes from Tukey's HSD test at $\alpha = 0.001$ are a, c, bc, b, respectively). Mean \log_{10} CFU values are given for each genotype inside the bar. Error bars represent SEM. Experiments shown in (a-d) were performed alongside each other. (e) Heatmap of mean expression values for genes associated with selected GO terms in indicated mutants relative to Col after syringe infiltration of Pst avrRps4 $(OD_{600} = 0.005)$. Shown GO terms were differentially expressed in one of the genotypes relative to Col $(|log_2FC| \ge 0.58 \text{ or } 1.5 \text{ times}, t\text{-test at FDR} < 0.05,$ asterisk shows where the GO terms are on the heatmap). Numbers next to the names of GO terms are the number of TPR1-bound genes vs the total number of genes in the corresponding GO term. The tpr1 and t3 mutants displayed a significant increase in the expression of genes from defense-related GO terms at 24 h (black arrow), for example, 'systemic acquired resistance' (SAR) and 'response to bacterium'. The '0 h' time point refers to c. 5 min after the infiltration. (f, g) Electrolyte leakage in Arabidopsis plants of indicated genotypes in response to nonvirulent Pseudomonas fluorescens bacteria Pf0-1 equipped with type III secretion system (T3SS) and expressing (f) or not (g) the avrRps4 effector (OD₆₀₀ = 0.2). The t3 mutant displayed increased electrolyte leakage at 24 h postinfection with these strains (Tukey's HSD, $\alpha = 0.001$; n = 16 from four independent experiments). Smaller differences were also detected at 6 hpi in tpr1 and t3 treated with Pf0-1 (T3SS, P < 0.05) but not at 0 hpi (ANOVA, P > 0.05). (h) The differential electrolyte leakage response in t3 is bacteria-triggered since infiltration of 10 mM MgCl₂ (infiltration medium, IM) gave similar conductivity levels in Col-0 and t3 at 24 h (ANOVA, P > 0.05). ns, not significant; CFU, colony-forming units. Elements of boxplots and matching statistics: first quartile – minima, third quartile – maxima, median – central line; whiskers extend to the minimum and maximum values but not further than 1.5 interquartile range from the respective minima or maxima of the boxplot. Datapoints with the same color were recorded in one independent experiment. Non-overlapping lowercase letter combinations above the bars and boxplots indicate statistically significant differences between the samples (Tukey's HSD test at $\alpha = 0.001$).

unknown function, *Plastid transcriptionally active 17 (PTAC17*, AT1G80480). We concluded that TPL/TPRs are largely dispensable for the transcriptional mobilization of defense in

TNL^{RRS1-RPS4}-mediated ETI to *Pst* bacteria, probably reflecting their limited contribution to bacterial growth restriction under our experimental conditions (Fig. 3a–d).

We applied a more sensitive analysis to test whether functionally coherent gene groups rather than individual genes are differentially expressed in tpr1 and t3 immune responses. For this, we used GO-based grouping of genes considering only GOs with 50-2000 genes. 76 of 350 tested GO-based gene sets were differentially expressed in one of the mutant lines (eds1, tpr1, and t3) at 0, 8, or 24 h relative to Col, as shown in heatmaps (Figs 3e, S6a) and Table S13 ($|\log_2 FC| \ge 0.5$, FDR ≤ 0.01). At 0 hpi (c. 5 min after Pst avrRps4 infiltration), eds1, tpr1, and t3 had reduced expression of genes with GO terms 'systemic acquired resistance' and 'response to bacterium' (Figs 3e, S6a,b), probably reflecting defects in a general stress response to leaf infiltration (Fig. S6c; Bhandari et al., 2019; Van Moerkercke et al., 2019). At 8 hpi, tpr1 and t3 mutants were more similar to Col than eds1 (Fig. S6a,b), underscoring the dispensability of TPL/TPRs for early transcriptional mobilization of resistance (Fig. 3e; Ding et al., 2020). Strikingly, at 24 hpi gene sets corresponding to GO terms 'systemic acquired resistance' and 'response to bacterium' had elevated expression in tpr1 and t3 mutants compared with Col (for the whole GO term, mean $log_2FC = 0.26-0.30$, or c. 1.2 times, FDR < 0.05; Figs 3e, S6a,b). Eight other GO terms such as 'response to fungus', 'leaf senescence', 'proteasome complex', and 'exocyst' also showed upregulation in tpr1 and t3 mutants (for the whole GO term, mean $log_2FC \ge 0.29$ or ≥ 1.95 times in t3, FDR < 0.05, full data in Table S13). These results suggest that TPL/TPRs contribute to the repression of defense gene expression after the initial wave of transcriptional elevation in a TNL^{RRS1-RPS4} ETI response. The transcriptome profiles of *tpr1* and *t3* in TNL^{RRS1-RPS4} ETI suggest a complex role of TPL/ TPRs in immunity depending on the stage of the immune response.

The *tpr1 tpl tpr4* mutant displays enhanced bacteria-triggered electrolyte leakage

Because several GO terms associated with immune responses were weakly upregulated in tpr1 and t3 at 24 hpi with Pst avrRps4 (Fig. S6a; Table S13), we tested whether TPL/TPRs help to restrict activated immune responses. In TNL RRS1-RPS4 ETI, host cell death measured as electrolyte leakage can be uncoupled from bacterial growth restriction (Heidrich et al., 2011; Lapin et al., 2019; Saile et al., 2020). We quantified electrolyte leakage in tpr1 and t3 mutants after infiltration of the type III secretion system (T3SS) equipped Pseudomonas fluorescens (Pf) 0-1 strain delivering avrRps4. At 24 h after Pf0-1 avrRps4 infiltration, conductivity was higher in Col than eds1, consistent with EDS1 being essential for TNL-triggered cell death (Heidrich et al., 2011; Lapin et al., 2019; Saile et al., 2020; Fig. 3f). While tpr1 plants behaved similarly to Col, the t3 mutant had increased conductivity at 24 hpi compared with Col (Fig. 3f). The same Arabidopsis lines were infiltrated with Pf0-1 empty vector (EV) that elicits strong PTI (Sohn et al., 2014; Saile et al., 2020). T3SS-equipped Pf0-1 (EV) also increased electrolyte leakage in the t3 mutant at 24 and weakly at 6 hpi compared with Col plants (Fig. 3g). No differences in electrolyte leakage were found between Col and t3 under mock treatment conditions (Fig. 3h).

These observations suggest that the *t3* mutant is defective in limiting bacteria-triggered cell death or damage.

We generated three independent stable homozygous lines expressing *pTPR1:TPR1-GFP* in the *t3* background. None of them displayed *TPR1* Col-like growth retardation or high TPR1-GFP protein accumulation (Fig. 4a,b). Despite having different TPR1-GFP amounts, these lines displayed reduced electrolyte leakage similar to wild-type Col and lower than *t3* (L2 and L3) or at the intermediate level (L1) at 24 hpi *Pf*0-1 EV infiltration (Fig. 4c). These data suggest a role of TPR1 in limiting bacteria-triggered electrolyte leakage. We propose that one potentially important and hitherto unknown role of TPR1 and other TPL/TPRs is to prevent an over-reaction of host tissues to pathogen infection.

TPL/TPRs reduce physiological damage associated with prolonged immunity

Genes with GO terms related to chloroplast functions were more strongly downregulated in tpr1 and t3 relative to Col at 24 hpi with Pst avrRps4 (Table S13). Also, TPR1-GFP was enriched at upstream sequences of *LHCA2* (light-harvesting complex gene 2) and nine other genes from these GO terms in TPR1 Col (Fig. 5a; Table \$13). Therefore, we hypothesized that TPR1 and other TPL/TPRs regulate photosynthetic performance during an immune response. We tested chlorophyll content and photosynthetic efficiency in healthy and bacteria-infected plants by quantifying photosystem II (PSII) fluorescence. While alterations in the operating PSII efficiency (\phiPSII) are measurable during shortterm stress, a drop in the maximum quantum yield of PSII $(F_{\rm v}/F_{\rm m})$ reflects more acute damage to PSII, and is observed under prolonged stress conditions (Baker, 2008). The tpr1 and t3 mutants were infiltrated with Pst bacteria ($OD_{600} = 0.005$) or mock solution (10 mM MgCl₂) alongside Col and eds1, which shows enhanced disease symptoms with Pst. Total chlorophyll content was reduced to a similar extent in Col, tpr1, and t3 upon Pst infection (Fig. 5b). A reduction in φ PSII and F_v/F_m values was minimal in infected Col leaves over the course of 3 d, indicating that these plants effectively balance bacterial growth restriction and PSII activity (Fig. 5c,d; purple line). By contrast, tpr1, t3, and eds1 had decreased ϕ PSII and F_v/F_m over 3 d following Pst infection (Fig. 5b,c; orange line – tpr1, blue line – t3). Reduced φ PSII and F_v/F_m in t3 and eds1 correlated with greater loss of fresh weight at 3 d after the Pst infiltration $(OD_{600} = 0.005, Fig. 5e)$. These observations suggest a role of TPL/TPRs in maintaining crucial photosynthetic functions in Pst-infected leaves, although confounding effects of the increased bacterial growth in t3 (Fig. 3a–d) cannot be excluded.

We used a pathogen-free system to test the role of TPL/TPRs in maintaining photosystem II functions upon activation of the immune system. PAMP nlp24 is recognized by the cell surface-localized receptor-like protein RLP23 (Oome *et al.*, 2014; Albert *et al.*, 2015; Seidl & Van den Ackerveken, 2019). RLP23 signals via the EDS1/PAD4 heterodimer in *Arabidopsis* (Pruitt *et al.*, 2021; Tian *et al.*, 2021). Although treatment with nlp24 did not alter F_v/F_m at 3 d (Fig. 5f), the operating PSII performance (ϕ PSII) was reduced in Col in an *EDS1*-dependent

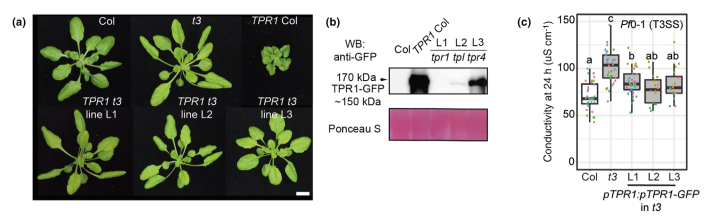


Fig. 4 *Arabidopsis* TPR1 counteracts electrolyte leakage triggered by the T3SS-equipped *Pf*0-1 bacteria. (a) Representative photographs of rosettes of 5–6-wk-old plants from three independent homozygous complementation lines expressing *pTPR1:TPR1-GFP* in *tpr1 tpl tpr4* (*t3*). *TPR1* Col is shown for comparison. The complementation lines do not show dwarfism in contrast to *TPR1* Col with the constitutive defense signaling. Scale bar = 1 cm. (b) Steady-state levels of TPR1-GFP in lines from (a), determined via Western blot analysis. Total protein extracts were probed with α-GFP antibodies. Ponceau S staining was used to control loading. The experiment was repeated two times with similar results. (c) Electrolyte leakage in the complementation lines from (a) and control lines Col and *t3* at 24 h after the *Pf*0-1 T3SS (OD₆₀₀ = 0.2) infiltration. The complementation lines L1–L3 show a level of the electrolyte leakage comparable to Col (Tukey's HSD α = 0.001; different colors of data points correspond to independent experiments, n = 12-24 from three or six independent experiments). Elements of boxplots and matching statistics: first quartile – minima, third quartile – maxima, median – central line; whiskers extend to the minimum and maximum values but not further than 1.5 interquartile range from the respective minima or maxima of the boxplot. Datapoints with the same color were recorded in one independent experiment. Non-overlapping lowercase letter combinations above boxplots indicate statistically significant differences between the samples (Tukey's HSD α = 0.001).

manner (Fig. 5g). The nlp24-triggered φPSII reduction was stronger in *t3* than in Col or the *t3 pTPR1:TPR1-GFP* L3 complementation line (Fig. 5g). This difference cannot be explained by increased perception of nlp24 in *t3* since *t3* was less sensitive to this PAMP than Col as measured in reactive oxygen species (ROS) burst assays (Fig. 5h,i). These results demonstrate that TPR1 and likely other TPL/TPRs contribute to maintaining PSII function in *EDS1*-dependent immunity.

We used the bacterial PAMP flg22 or phytocytokine Atpep1 (pep1 hereafter) to evaluate the effect of TPL/TPRs on the growth of immune-triggered plants in a pathogen-free system (Fig. 6). Root growth inhibition (RGI) was similar in Col, *tpr1*, and t3 mutants in the presence of 100 nM flg22 (Fig. 6a,b), but it was more pronounced in the t3 mutant on medium supplemented with 50 or 200 nM pep1 (Fig. 6c-e). RGI hypersensitivity of t3 seedlings to pep1 was rescued in the TPR1-GFP complementation lines (Fig. 6c-e). Perception of pep1 was not altered in t3 compared with Col at the level of pep1-induced phosphorylation of mitogen-activated protein kinase 3 and 6 (MPK3 and MPK6; Fig. 6f). Also, t3 seedlings showed Col-like induction of PTI marker genes FRK1 (FLG22-induced receptorlike kinase 1) and WRKY30 at 1 h after the flg22 or pep1 treatment (Fig. 6g). Hence, TPR1 and most likely other TPL/TPRs reduce RGI in phytocytokine-stimulated sterile seedlings. Taken together, the data suggest that Arabidopsis TPR1 and other TPL/ TPRs limit physiological and growth penalties associated with induced immunity.

Discussion

Timely activation of immune responses is essential for plant resistance to pathogens. How activated defenses are subsequently

restricted to prevent damaging over-stimulation of tissues is less clear. Here, we present evidence that TPR1 and probably other TPL/TPR family transcriptional corepressors contribute to limiting host physiological damage and growth inhibition associated with induced immunity.

We obtained ChIP-Seq chromatin association profiles for Arabidopsis TPR1 with or without constitutive EDS1-dependent defense. TPR1-GFP bound primarily to upstream regions of c. 1400 genes, and c. 10% of these genes showed enhanced TPR1-GFP binding when EDS1-dependent immunity signaling was active (Fig. 2). TPR1-GFP association with regions immediately upstream of the transcription start site (TSS) is consistent with TPR1 interaction with TFs (Szemenyei et al., 2008; Causier et al., 2012) and with the location of predicted TF-binding sites close to the TSS (Yu et al., 2016). Although TPR1-bound genes are strongly enriched for ChIP signals of MYC2 (Van Moerkercke et al., 2019; Wang et al., 2019; Zander et al., 2020), WRKYs (Birkenbihl et al., 2018), and SARD1 (Sun et al., 2015) TFs (Fig. 2g), there is so far no reported evidence for complex formation between these TFs and TPR1 in plant tissues. TPR1-GFP-bound genes are enriched for GO terms associated with defense and control of growth and development (Tables S6, S8). Also, binding of TPR1 to chromatin correlated with transcriptional activation (Fig. 2). However, we cannot draw a conclusion about causality between TPR1 binding and the transcriptional output. Time series ChIP-Seq and RNA-Seq experiments with complementation lines would be useful to address this question. Because of redundancy between TPL/TPR members, we think that the processed input-normalized TPR1-GFP chromatin enrichment profiles for TPR1 Col and TPR1 eds1 provided here could help researchers identify chromatin interactions for TPL/ TPRs (see Data availability section).

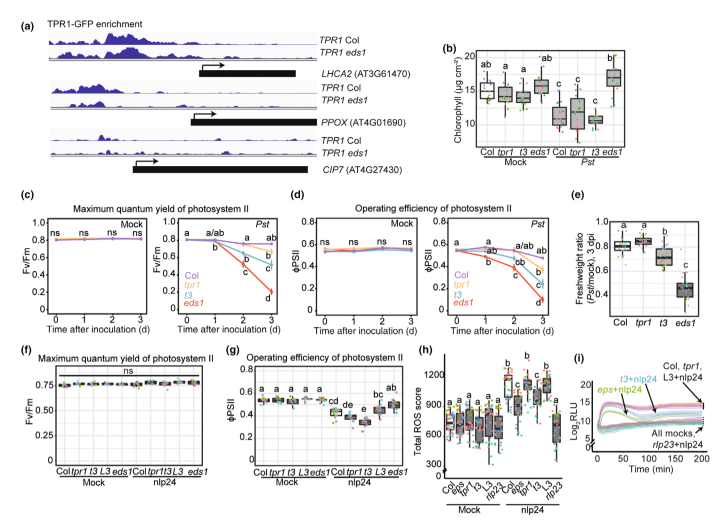


Fig. 5 TPR1 limits adverse effects of nlp24-activated PTI on Arabidopsis PSII performance. (a) Genes with chloroplast functions are direct binding targets of TPR1. Input-normalized ChIP-Seq profiles show that TPR1-GFP binds in TPR1 Col and TPR1 eds1 at the upstream sequences of LHCA2 (AT3G61470), PPOX (AT4G01690), and CIP7 (AT4G27430) involved in photosynthesis or chlorophyll biosynthesis. (b) tpr1 and t3 mutants display the Col-like chlorophyll reduction after Pst infection. Plants of indicated genotypes were syringe-infiltrated with Pst (OD₆₀₀ = 0.005) and total (a + b) chlorophyll content was determined at 3d post-inoculation (dpi) (Tukey's HSD $\alpha = 0.05$; n = 15 from three independent experiments). (c, d) Maximum quantum yield of PSII (F_v/F_m) (c) and operating PSII efficiency (φPSII) (d) in indicated genotypes over the 3-d time course after syringe infiltration of Pst (OD₆₀₀ = 0.005; left panels). Compared with Col, the t3 mutant shows significantly reduced F_v/F_m at 3 d after infection with Pst but not in the mock-treated samples (Tukey's HSD $\alpha = 0.05$; n = 9-12 from three independent experiments). Error bars represent SEM. (e) Fresh weight reduction in leaves inoculated with Pst (OD₆₀₀ = 0.005) compared with mock-treated leaves in indicated genotypes 3 dpi (Tukey's HSD $\alpha = 0.05$; n = 20-24 from four independent experiments). (f, g) F_v/F_m (f) and φ PSII (g) in indicated genotypes 3 dpi after syringe infiltration of 5 μ M nlp24 or mock (10 mM MgCl₂) control (Tukey's HSD α = 0.05; n = 10–14 from four independent experiments). φPSII drops stronger in the t3 mutant but this is restored to Col levels in the complementation line L3 (pTPR1:TPR1-GFP L3; t3 background). (h, i) Reactive oxygen species burst in the indicated genotypes in the presence of nlp24 (2 µM). (h) Total ROS score is a sum of log₂transformed relative luminescence units (RLU) recorded over the time course 200 min. ROS burst is lower in the t3 and eds1-12 pad4-1 sag101-3 (eps) mutant compared with Col and the L3 complementation line. (i) The difference between Col and t3 (blue ribbon) is visible after c. 1 h after exposure to nlp24 (Tukey's HSD $\alpha = 0.05$; n = 27-30 from five independent experiments). Shadowing around lines corresponds to one SEM. ns, not significant. Elements of boxplots and matching population statistics: first quartile - minima, third quartile - maxima, median - central line; whiskers extend to the minimum and maximum values but not further than 1.5 interquartile range from the respective minima or maxima of the boxplot. Datapoints with the same color were recorded in one independent experiment. Non-overlapping lowercase letter combinations in (b-e, g-i) indicate that the samples have statistically significant differences (Tukey's HSD $\alpha = 0.05$).

Our RNA-Seq gene expression data do not support the role of TPR1 and other TPL/TPRs in *EDS1*-dependent transcriptional host defense mobilization at 8 or 24 h after bacterial inoculation despite compromised resistance of the *t3* mutant to different *Pst* strains (Fig. 3a–d). How then do TPL/TPRs positively regulate *Arabidopsis* defense against bacteria? An earlier report showed

that TPR1 and other TPL/TPRs contribute to PAMP-triggered ROS burst from the surface receptor FLS2 (Navarrete et al., 2021) that is largely EDS1-independent (Pruitt et al., 2021; Tian et al., 2021). We find that nlp24-triggered RLP23 and EDS1 family-dependent ROS burst is reduced in the t3 mutant but it recovered in the complementation lines

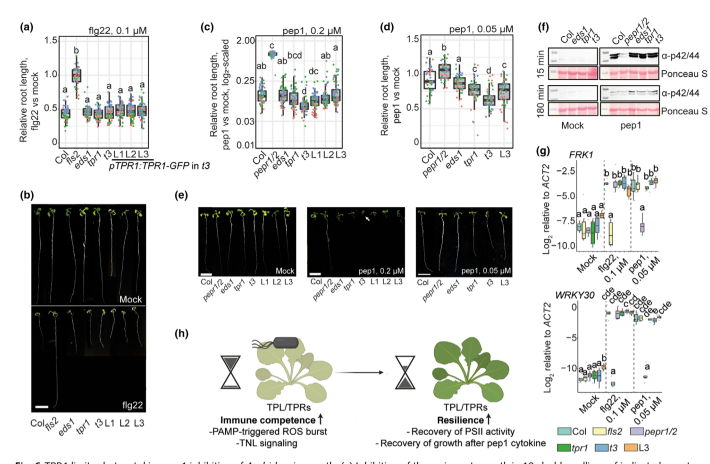


Fig. 6 TPR1 limits phytocytokine pep1 inhibition of Arabidopsis growth. (a) Inhibition of the main root growth in 10-d-old seedlings of indicated genotypes grown on liquid 0.5× MS supplemented with flg22 (100 nM). The eds1, tpr1, t3 mutants as well as the pTPR1:TPR1-GFP complementation lines (L1–L3) showed the Col-like root growth inhibition, while the flg22 receptor mutant fls2 was insensitive to the treatment (Tukey's HSD $\alpha = 0.05$; n = 58 from three independent experiments). The main root length in flg22-treated plants was normalized to the respective mean in mock-treated seedlings. (b) Photographs of representative seedlings measured in (a). Bar, 1 cm. (c, d) Boxplot representation of root growth ratio after exposure of 10-d-old seedlings of indicated genotypes on $0.5 \times$ liquid MS medium with $0.2 \,\mu$ M (c) or $0.05 \,\mu$ M (d) pep1 relative to the mock (mQ water) treatment (Tukey's HSD $\alpha = 0.05$; n = 58 and n = 48 from three independent experiments in c and d, respectively). Y-axis in (c) is log_2 -scaled. The t3 mutant is hypersensitive to pep1 at the level of root growth inhibition, but this is recovered in the L3 complementation line. (e) Representative photographs of seedlings from (c) and (d). Bar, 1 cm. (f) MPK3 and MPK6 phosphorylation assessed via Western blot analysis with α -p42/44 antibodies in indicated genotypes at 15 and 180 min after mock (mQ water) or pep1 (0.2 μM) treatment. The t3 mutant showed Col level of MPK3 and MPK6 phosphorylation. The experiment was repeated three times with similar results. In (a-f), eds1 is eds1-12. (g) Induction of PTI marker genes FRK1 (FLG22-induced receptor-like kinase 1) and WRKY30 in seedlings of indicated genotypes after exposure to flg22 (0.1 μM), pep1 (0.05 μM) or mock (mQ) for 1 h. Seedlings were grown on liquid 0.5 × MS for 11 d (red datapoints) or 12 d (blue datapoints). The tpr1, t3 mutants and the complementation line L3 responded similarly (Tukey's HSD $\alpha = 0.05$; n = 6 from two independent experiments). (h) Model of the function of TPR1 and other TPL/TPRs in immune-triggered Arabidopsis leaves. TPL/TPRs promote immune activation in the initial response (e.g. ROS production) but help to maintain PSII activity and growth at later stages of the immune responses. The picture was created with BioRender.com. Elements of boxplots and matching population statistics: first quartile – minima, third quartile – maxima, median –central line; whiskers extend to the minimum and maximum values but not further than 1.5 interquartile range from the respective minima or maxima of the boxplot. Datapoints with the same color were recorded in one independent experiment. Non-overlapping letter combinations above the boxplots in (a, c, d, g) show statistically significant differences between the samples (Tukey's HSD $\alpha = 0.05$).

(Fig. 5h,i). Thus, TPR1 and probably other TPL/TPRs contribute to a PTI-associated ROS burst. This defect in PTI might explain weakly reduced expression of gene sets from GO terms linked to immunity in *tpr1* and *t3* mutants after *Pst avrRps4* infiltration (Fig. 3e; *c.* 5 min). Although TPR1 contributes to TNL snc1 signaling (Zhu *et al.*, 2010; Zhang *et al.*, 2019; Navarrete *et al.*, 2021), which in turn requires the EDS1 family, it is still unclear whether a TPL/TPRs contribution to PAMP ROS burst occurs via TIR domain signaling. In addition, TPR1 was proposed to induce defenses by repressing negative regulators of

resistance (Zhu *et al.*, 2010). Consistent with this view, TPR1 is enriched at promoters of genes that are repressed during TNL^{RRS1-RPS4} ETI (Fig. S3; Bartsch *et al.*, 2006; Zhu *et al.*, 2010), and TPR1 can repress *DND1/CNGC2* and *DND2/CNGC4* promoter activity (Niu *et al.*, 2019).

The *t3* mutant over-responded at the level of defense-associated gene expression at 24 hpi with *Pst avrRps4* bacteria and bacteria-triggered electrolyte leakage (Fig. 3). Also, we identified a contribution of TPR1 and other TPL/TPRs to supporting photosystem II activity in leaves infected with *Pst* or infiltrated

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with PAMP nlp24 and in maintaining seedling growth in response to pep1 (Figs 5, 6). Hence, a model emerges in which *Arabidopsis* TPR1 and other TPL/TPRs transcriptional corepressors promote initial immune activation (Figs 3e, 5i) but subsequently mitigate adverse effects of activated immunity signaling on host physiology and growth (Fig. 6h). Interestingly, TPR2 and TPR3 likely work as negative regulators of TNL snc1-conditioned autoimmunity and shoot growth retardation (Garner *et al.*, 2021). Thus, both clades of the TPL/TPR family, TPR1/TPL/TPR4 and TPR2/TPR3, can help to reduce damaging effects of activated immunity in *Arabidopsis*.

Timely downregulation of defense signaling is relevant because prolonged pathogen infection and plant immune activation often lead to reduced photosynthetic activity and biomass (Walters, 2015a,b). Pathogen-free induction of SA and JA signaling is associated with reduced expression of genes involved in photosynthesis (Hickman et al., 2017, 2019). Despite the identification of multiple genes affecting the balance of growth and defense (Huot et al., 2014; Bruessow et al., 2021), knowledge of how plants turn down transcriptional defenses and regain physiological homeostasis is fragmentary. Cytoplasmic condensates of the SA receptor NPR1 were reported to be responsible for the ubiquitination of ETI cell death-promoting WRKY TFs to limit their activities (Zavaliev et al., 2020). Also, the SA receptor NPR4 suppresses Arabidopsis WRKY70 promoter activity (Ding et al., 2018). Our study identifies Arabidopsis transcriptional corepressor TPR1 as a factor that prevents overshooting of an immune response and therefore as a potential contributor to plant stress-fitness balance.

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Competing interests

None declared.

Author contributions

TG, DL, FL, and JEP designed experiments. TG, DL, and FL performed experiments. TG, DL, and JEP analyzed all data. BK, LC, and MB analyzed ChIP-Seq and RNA-Seq. JB and DL generated and characterized complementation lines. DL prepared

the Github repository and materials for access to processed ChIP-Seq data. TG, DL, and JEP wrote the manuscript with input from all authors. TG and DL contributed equally to this work.

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Data availability

RNA-Seq and ChIP-Seq data from this article are deposited in the National Center for Biotechnology Information Gene Expression Omnibus (GEO) database with accession nos. GSE149316, GSE154652, and GSE154774. Bigwig, BAM, and BAI files of TPR1 ChIP-Seq for visualization in IGV browser are also available through the Max Planck Digital Library collection (MPDL; https://edmond.mpdl.mpg.de/imeji/collection/U6N5zIOIWgjjMZCu). Scripts for preparing metaplots in R environment on a personal computer (c. 8G RAM) are on GitHub (https://github.com/rittersporn/TPR1_metaplots_Griebel_Lapin_etal_2021).

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Supporting Information

Additional Supporting Information may be found online in the Supporting Information section at the end of the article.

- **Fig. S1** Autoimmunity of *TPR1* Col is *EDS1*-dependent.
- **Fig. S2** Input-normalized ChIP-Seq profiles of TPR1-GFP in *TPR1* Col and *TPR1 eds1* at selected genes showing differential TPR1-GFP binding and expression in *TPR1* Col and *TPR1 eds1*.
- **Fig. S3** Input-normalized ChIP-Seq profiles of TPR1-GFP in *TPR1* Col and *TPR1* eds1 at the indicated genes bound by hemagglutinin tagged TPR1 in autoimmune *TPR1-HA* Col line in Zhu et al. (2010).
- **Fig. S4** Input-normalized ChIP-Seq profiles of TPR1-GFP in *TPR1* Col and *TPR1 eds1* at genes known as targets of TPL.
- **Fig. S5** MYC, WRKYs, and SARD1 chromatin binding events are enriched at the TPR1-bound loci.
- **Fig. S6** The *tpr1 tpl tpr4* mutant showed enhanced defense transcriptional reprogramming at 24 hpi with *Pst avrRps4*.
- Methods S1 Methods (continued).
- Table S1 Oligonucleotides used in this study.
- **Table S2** Results of differential gene expression analysis for Col-0 (Col), *TPR1* Col, *TPR1 eds1*, and *TPR1 sid2*.

- **Table S3** Placement of genes differentially expressed between Col, *TPR1* Col, *TPR1 eds1*, and *TPR1 sid2* in different clusters after hierarchical clustering of their expression profiles.
- **Table S4** Results of the Gene Ontology (GO) term enrichment analysis on gene clusters in Table S2.
- **Table S5** Annotation of TPR1-GFP peaks called in *TPR1* Col ChIP-Seq.
- **Table S6** Results of the GO term enrichment analysis on genes associated with TPR1 peaks in *TPR1* Col.
- **Table S7** Annotation of TPR1-GFP peaks called in *TPR1 eds1* ChIP-Seq.
- **Table S8** Results of the GO term enrichment analysis on genes associated with TPR1 peaks in *TPR1 eds1*.
- **Table S9** Differential TPR1 enrichment at the chromatin regions in *TPR1* Col and *TPR1 eds1*.
- **Table S10** TPR1-binding categories according to Fig. 2(e) with Gene IDs
- **Table S11** Overlap between DEGs in different datasets and TPR1 targets from Fig. 2(e).
- **Table S12** Results of the differential gene expression analysis for Col-0 (Col), Col-0 *eds1-2* (*eds1*), *tpr1*, and *tpr1 tpl tpr4* (*t3*) at 0, 8, and 24 h after infection with *Pst avrRps4*.
- **Table S13** Results of the gene set expression analysis for Col, *eds1*, *tpr1*, and *t3* at 0, 8, and 24 h, respectively, after infection with *Pst avrRps4*.

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