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# Downy Mildew effector HaRxL21 interacts with the transcriptional repressor TOPLESS to promote pathogen susceptibility

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# Abstract

*Hyaloperonospora arabidopsidis (Hpa)* is an oomycete pathogen causing Arabidopsis downy mildew. Effector proteins secreted from the pathogen into the plant play key roles in promoting infection by suppressing plant immunity and manipulating the host to the pathogen's advantage. One class of oomycete effectors share a conserved 'RxLR' motif critical for their translocation into the host cell. Here we characterize the interaction between an RxLR effector, HaRxL21 (RxL21), and the Arabidopsis transcriptional co-repressor Topless (TPL). We establish that RxL21 and TPL interact via an EAR motif at the C-terminus of the effector, mimicking the host plant mechanism for recruiting TPL to sites of transcriptional repression. We show that this motif, and hence interaction with TPL, is necessary for the virulence function of the effector. Furthermore, we provide evidence that RxL21 uses the interaction with TPL, and its close relative TPL-related 1, to repress plant immunity and enhance host susceptibility to both biotrophic and necrotrophic pathogens.

# Author summary

Disease in plants depends on the outcome of interaction between the pathogen and the host plant. Pathogens use effector proteins to aid colonization by suppression of the host immune system. Identification of effector targets can pinpoint key components of plant defence in addition to aiding our understanding of how pathogens manipulate their hosts. Here, we see a downy mildew effector mimicking a plant transcriptional repression motif and interacting with a transcriptional co-repressor in order to manipulate the plant immunity transcriptional response and susceptibility to both a biotrophic and a necrotrophic pathogen.

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#### Introduction

Plants are constantly under attack by pathogenic microbes. In most cases, the microbe is not able to cause disease on a particular plant species due to pre-formed barriers to infection and/ or the ability of the plant to recognize conserved molecular motifs, known as microbe (or pathogen) associated molecular patterns (MAMPs or PAMPs), and activate additional defence responses [1,2]. Signaling pathways activated downstream of MAMP recognition (MAMP or PAMP-triggered immunity; PTI) by pattern recognition receptors (PRRs) result in production of reactive oxygen species, hormone biosynthesis, callose deposition in the cell wall and large-scale transcriptional reprogramming within the plant [3,4]. Many pathogens use effector proteins to suppress or evade these host immune responses and/or adapt host physiology to aid infection [5]. For example, effectors may target PTI signalling [6–9] or manipulate stomatal opening [10]. Manipulation of the host plant by the pathogen also occurs through alteration of host transcription; it is well documented that transcription activator-like effector (TALe) proteins from the *Xanthomonas* genus of plant pathogenic bacteria bind and activate host promoters [11]. Large-scale changes in host transcription brought about by effectors have also been shown during infection by *Pseudomonas syringae* py. *tomato* DC3000 (*Pst*) [12,13].

Alignment of known effector proteins from plant pathogenic oomycetes including *Hyaloperonospora arabidopsidis* (*Hpa*) revealed a consensus sequence RxLR (arginine, any amino acid, leucine, arginine) downstream from a signal peptide [14], facilitating identification and subsequent characterization of candidate effector proteins from the *Hpa* genome [15,16]. Potential Arabidopsis protein targets of *Hpa* effectors have been identified using yeast-2-hybrid (Y2H) with 122 different Arabidopsis proteins targeted by 53 *Hpa* effectors [17,18]. One effector from *Hpa*, HaRxL21 (RxL21) was found to interact with the Arabidopsis transcriptional corepressor TOPLESS (TPL) [17].

In Arabidopsis, the TPL family consists of five members, TPL and four TPL-related (TPR) proteins. TPR1 is the most similar to TPL sharing 95% similarity at the amino acid level [19–21]. TPL and TPRs have been shown by Y2H to interact with several regulators of hormone pathways known to be involved in plant defence against pathogens [22], abiotic stress and in development [23]. Moreover, involvement of TPL/TPRs in regulation of resistance protein signalling [20,24] and jasmonate signalling [25] has been shown. We speculate that TPL is a key target for manipulation by pathogens.

The TPL family is highly conserved in land plants [26,27] and shows structural similarity to the *Drosophila* protein Groucho, in addition to human transducin-like Enhancer of split and transducin (beta)-like 1 proteins [28–30]. TPL family members link transcription factors (TFs) to chromatin remodelling complexes; the corepressors interact with TFs and recruit chromatin remodelling factors such as histone deacetylases [20,21,31]. For example, the Arabidopsis regulators of root hair development GIR1 and GIR2 have been shown to promote histone hypoace-tylation via their interaction with TPL [32]. The current understanding is, therefore, that TPL/TPR recruitment to a gene results in reversible transcriptional repression via condensed DNA structure that is less accessible to transcription initiation complexes.

At the very N-terminus TPL harbours a LIS1 homology domain (LisH) followed by a C-terminal to LisH (CTLH) domain. The latter is needed for the interaction with proteins containing an Ethylene-responsive element Binding Factor-associated Amphiphilic Repression (EAR) motif [33,34]. The EAR motif was first identified as the conserved sequence  $^{L}/_{F}DLN^{L}/_{F}$  (x)P in class II Ethylene Response Factor genes which function as transcriptional repressors [35] and subsequently further characterised as LxLxL or DLNxxP [36]. EAR-mediated protein-protein interaction with TPL is often required for transcriptional repression. For example, the transcriptional regulator IAA12 requires interaction between its EAR motif and the CTLH domain of TPL for its repressive activity in low auxin conditions [33,34]. Repression of jasmonate signalling relies on interaction between TPL and the EAR motif-containing Novel Interactor of JAZ (NINJA) [25,37]. Interaction with TPL and transcriptional repressor activity of NINJA were abolished when Leu residues in the EAR motif were mutated to Ala [25].

EAR motifs have now been identified in many plant proteins involved in development, stress and defence [38–40]. So far, a few pathogen effectors mimicking this EAR motif have been described although little is known about subsequent corepressor recruitment. For example, the *Xanthomonas campestris* type III effector XopD has been found to contain two tandemly repeated EAR motifs downstream of a DNA binding domain which are required for XopD-dependent virulence in tomato [41,42]. Effectors from the XopD superfamily that contain conserved EAR motifs have been found in *Xanthomonas, Acidovorax* and *Pseudomonas* species [41]. In addition, a conserved EAR motif is required for virulence in the *Ralstonia solanacearum* effector PopP2 although again no interaction with known Arabidopsis corepressors has been identified so far [43]. There are however, examples of pathogen effectors interacting with TPL family members including interaction of the *Melampsora larici-populina* effector MLP124017 with TPR4 [44].

The aim of our work was to determine how RxL21 is manipulating Arabidopsis to promote infection by *Hpa*. We demonstrate that RxL21 interacts *in planta* with the Arabidopsis corepressors TPL and TPR1 via an EAR motif and that this interaction is essential for RxL21 virulence activity against both Hpa and a necrotrophic plant pathogen. We find there is cooccurrence of RxL21 and TPR1 binding sites on promoter regions of a set of TPR1-repressed defence-related genes, suggesting that RxL21 virulence function involves perturbation of TPL/ TPR1 transcriptional repression during mobilization of host immunity.

#### Results

# **RxL21** is conserved across multiple *Hpa* isolates and contains a C-terminal EAR motif

RxL21 is a 45 kDa effector protein identified from the genome of *Hpa* [16]. It contains a 'RLLR-DEER' motif at the N-terminus and an EAR (LxLxL) motif (amino acid sequence LMLTL) at the C-terminus. Alleles of *RxL21* have been found in *Hpa* isolates Cala2, Emco5, Emoy2, Emwa1, Hind2, Maks9, Noks1 and Waco9 ([45] and BioProject PRJNA298674); alignment of their amino acid sequences is shown in S1 Fig. The signal peptide cleavage site was predicted to be between position 16 and 17 (SignalP-5.0). The RxLR-DEER motif is conserved across all alleles and with the exception of Noks1 (truncated due to Serine 197 changed to a Stop codon) the EAR motif at the C-terminus is also conserved between all aligned alleles. *RxL21* is expressed in conidiospores of virulent (Waco9) and avirulent (Emoy2) *Hpa* isolates during Col-0 infection, as well as in Waco9 1, 3 and 5 days post inoculation [46] consistent with it playing a role in virulence.

# RxL21 expression *in planta* causes enhanced susceptibility to both biotrophic and necrotrophic pathogens

Constitutive expression of RxL21 (from *Hpa* isolate Emoy2) in Arabidopsis has been shown to enhance growth of *Hpa* isolate Noco2 and a *Pst* isolate impaired in the suppression of early immune responses (*Pst* DC300  $\Delta$ avrPto/ $\Delta$ avrPtoB) [16]. In addition, RxL21 increased bacterial growth when delivered into Arabidopsis via the type-three secretion system of *Pst* DC3000-LUX [16]. Here, Arabidopsis plants overexpressing RxL21 [16] were screened for susceptibility to *Hpa* isolates Noks1 and Maks9, and the necrotrophic pathogen *Botrytis cinerea*.



**Fig 1. RxL21 expression** *in planta* **causes enhanced susceptibility to biotrophic and necrotrophic pathogens.** (A) Transgenic Arabidopsis expressing RxL21 under a 35S promoter (RxL21a/b) were challenged with *Hpa* isolates Noks1 and Maks9 and sporangiophores counted at 4 dpi. Col-0 WT and 35S::GUS (Col-0 background) were used as controls. (Noks1 n = 45, Maks9 n = 55). (B) RxL21a/b were challenged with *B. cinerea* and lesion area measured 72 h post infection (n = 24). Box plots show the median, upper and lower quartiles, whiskers show the upper and lower extremes of the data. Letters indicate significant difference using a Kruskal Wallis test with p < 0.05. Experiments were repeated with similar results.

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RxL21 lines were compared to both Col-0 wild type (WT) and lines expressing 35S::GUS (Col-0 GUS) which had been transformed and selected alongside the effector lines. Presence of the effector conferred enhanced susceptibility to both obligate biotroph *Hpa* isolates in two independent transgenic lines (RxL21a, b) compared to both Col-0 and Col-0 GUS, measured by total sporangiophores per seedling at 4 days post inoculation (dpi) (Fig 1A). RxL21 expression also resulted in increased lesion size caused by *B. cinerea* infection compared to controls (Fig 1B). Hence, it appears that the RxL21 effector is targeting a mechanism essential for a full defence response against both biotrophic and necrotrophic pathogens.

#### RxL21 interacts with the CTLH domain of TPL via its EAR motif

It has been previously reported that RxL21 interacts with TPL using the matrix yeast two hybrid technique (Y2H) [17]. To confirm that this interaction could occur in planta, we first determined that RxL21 and TPL co-localise to the nucleus when transiently expressed in Nicotiana benthamiana leaves (Fig 2A). We next investigated the specific interaction motifs of both partners, which have not been previously reported. Two EAR-mutation constructs  $(RxL21\Delta EAR1; \Delta 402-409 \text{ and } RxL21\Delta EAR2; \Delta 360-409)$  were used to determine whether the amino acids flanking the EAR motif are necessary for interaction with TPL (Fig 2B). As expected, full-length RxL21 showed a positive interaction with TPL via activation of histidine (GAL1::HIS3) and adenine (GAL2::ADE2) reporter genes. The screen was performed in both directions with RxL21 and TPL fused to both the activation domain (AD) or the DNA binding domain (DB). Selective plates containing 3AT (a competitive inhibitor of the HIS3 gene product) were used as an additional control for increased stringency. Binding affinity appeared to be influenced by the direction of cloning with activation of the adenine reporter gene (and high stringency activation of the histidine reporter) only detected when TPL was fused to the DB domain. Deletion of either the initial N terminal sequence ( $RxL21\Delta N$ ) or the RxLR-DEER motif (RxL21ARxLR) did not prevent interaction with TPL. However, deletion of the RxL21 EAR motif abolished the interaction with TPL (Fig 2B), as did deletion of the CTLH domain



Fig 2. The EAR motif of RxL21 interacts with the CTLH domain of TPL. (A) RxL21 co-localizes with TPL in the nucleus. GFP:RxL21 and RFP:TPL were expressed transiently in N. benthamiana. Scale bars are 10 µm. (B) TPL Interacts with the EAR domain of RxL21 in yeast. RxL21 was cloned without the signal peptide and lacking the EAR motif (RxL21 $\Delta$ EAR1;  $\Delta$ 402–409 and HaRxL21 $\Delta$ EAR2;  $\Delta$ 360–409), without the N-terminus (RxL21 $\Delta$ N;  $\Delta$ 1–36) and RxLR-DEER motif (RxL21ARxLR; Δ1-51) and combinations of the above. Amino acid locations of deletion constructs are indicated. Deleting the EAR motif at the C-terminus of RxL21 abolishes interaction with TPL by Y2H. Successful mating is indicated by growth on media lacking Leucine and Tryptophan (-LW). Growth on media additionally lacking Histidine (-LWH) indicates GAL1::HIS3 reporter gene activation and protein-protein interaction. 3AT is used to increase stringency in a concentration dependent manner. Growth on media additionally lacking Adenine (-LWHA) indicates activation of the GAL2::ADE2 reporter gene. AD; activation domain. DB; DNA binding domain. (C) Deleting the CTLH domain of TPL (TPL $\Delta$ CTLH;  $\Delta$ 25–91) abolishes interaction with RxL21 in yeast. Interaction (indicated by growth on-LWH +1mM 3AT) was observed between RxL21 and TPL but not between RxL21 and TPLACTLH. All Y2H was repeated at least twice with similar results. (D) TPL interacts with RxL21 and the interaction requires the EAR motif in planta. TPL:HA, TPLACTLH:HA together with RxL21 and RxL21 EAR motif mutants RxL21ΔEAR, RxL21-FMFTF and RxL21-AMATA (under control of an estradiol-inducible promoter and N-terminal Myc tag) were transiently expressed in N. benthamiana leaves. RxL21 expression was induced by 30μM β-estradiol 24 hr prior to harvesting, c-myc beads were used for immunoprecipitation (IP). HA antibody was used for TPL and TPLACTLH immunoblots (IB) and c-myc antibody was used to detect RxL21, RxL21AEAR and the respective EAR motif mutants.

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of TPL (Fig 2C). Hence, both the EAR motif of RxL21 and the CTLH domain of TPL are necessary for direct protein-protein interaction. The mechanism of interaction between the effector and its host target (TPL) is mimicking the mechanism by which plant proteins (such as NINJA [25]) interact with TPL and recruit it to a complex for transcriptional repression.

#### Leucine residues in the EAR motif are critical for interaction with TPL

The three Leucine (Leu/L) residues (Leu389, Leu391, Leu393) in the EAR motif of RxL21 were mutated individually, in pairs or all together to Phenylalanine (Phe/F), Alanine (Ala/A) and Isoleucine (Ile/I). Substitution to Phe was chosen because previously Leu to Phe mutation of Leu354, Leu356 and Leu358 in the Arabidopsis repressor SUPERMAN prevented corepressor activity [47]. In addition, the three Leu residues in RxL21 were mutated to Ala (as it lacks functional side chains) [48] and Ile because it is the most similar amino acid to Leu [49].

Mutagenesis of all three Leu residues to Phe, Ala or Ile was found to abolish interaction of RxL21 with TPL by Y2H (S2 Fig). Mutation of the 'x' residues within the EAR motif between the Leu residues (Methionine390 and Threonine392) had no effect on the interaction. Mutation of the first Leu residue (Leu389) or any pair of Leu residues in the EAR motif was sufficient to abolish RxL21-TPL interaction (S2A Fig). This analysis was repeated using TPL from *N. benthamiana* (*Nb*TPL). RxL21 interacted with *Nb*TPL and the same residues in the RxL21 EAR motif were necessary for interaction with *Nb*TPL as for interaction with *At*TPL (S2B Fig). These data are consistent with previous observations that mutation of any two Leucine residues within the DLELRL hexapeptide of SUPERMAN is sufficient to abolish repression of transcription [47].

#### **RxL21 and TPL interact** in planta

Bimolecular fluorescence complementation (Split YFP) was used to verify the interaction *in planta*. RxL21 was cloned into a vector with an N-terminal fused YFP fragment to minimise steric hindrance to the C-terminal EAR motif. *Agrobacterium tumefaciens* harbouring the N and C fragments of E-YFP fused to RxL21 and TPL were infiltrated into *N. benthamiana* and imaged using confocal microscopy. YFP fluorescence was detected in the nucleus, but no fluorescence was observed when RxL21ΔEAR was used instead of the full-length effector (S3 Fig).

The interaction between RxL21 and TPL was confirmed by co-immunoprecipitation (Co-IP) using transient expression in *N. benthamiana*. A Myc tag was fused to the N terminus of RxL21 (Myc-RxL21) and co-expressed with C-terminal HA-tagged TPL (TPL-HA). Immunoprecipitation of Myc-RxL21 using an anti-Myc antibody resulted in Co-IP of TPL-HA (Fig 2D) confirming direct protein-protein interaction occurs *in planta*. Furthermore, Myc-tagged RxL21ΔEAR was unable to pull down TPL, and TPLΔCTLH was not immunoprecipitated by Myc-RxL21, demonstrating that, as in Y2H, the EAR motif and CTLH domain are required for RxL21-TPL interaction. Variants of RxL21 in which the Leu residues in the EAR motif were mutated to Phe or Ala (RxL21-FMFTF or RxL21-AMATA) were also unable to pull down TPL, confirming our observations in yeast that the Leu residues in the EAR motif of RxL21 are necessary for interaction with TPL.

#### An EAR motif is not sufficient to mediate interaction with TPL

In a previous large-scale study, no other *Hpa* effectors were found to interact with TPL [17]. The amino acid sequence of 134 RxLR (RxL), 491 RxL-like (RxLL) and 20 Crinkler (CRN) candidate effector proteins in the *Hpa* genome [15,45] were searched for the presence of the LxLxL EAR motif. The predicted signal peptide cleavage site in the effectors was then

identified using SignalP [50] and any effectors containing an LxLxL motif within the predicted signal peptide were subsequently excluded from further analysis.

The LxLxL motif was found to be present in 16 RxLs and 35 RxLLs (S1 Table). Of these, 2 RxLs and 4 RxLLs were found to contain multiple LxLxL motifs. The position of the LxLxL motif was noted, since previous reports of the EAR motif in functionally characterized transcriptional repressors is often at the N or C terminus [25,51,52]. The LxLxL motif of 5 RxLs (including RxL21) and 7 RxLLs was found to be within 35 amino acids of the C-terminus of the protein. No RxLs or RxLLs were found to have a LxLxL motif at the N terminus. We compared LxLxL-containing effectors against available subcellular localization data [53] and found, in addition to RxL21, HaRxL48, HaRxL55, HaRxLL100 and HaRxLL470 were nuclear localised in *N. benthamiana* [53], suggesting a possible functional role for the EAR motif in these effectors in manipulating host transcription. Ten of the LxLxL-containing *Hpa* effectors (including three of the four nuclear-localised ones) were tested using Y2H. None of the effectors showed an interaction with TPL (S4 Fig) indicating that the presence of an EAR motif is not sufficient to mediate an interaction with TPL and that RxL21 may have a unique function.

## The EAR domain is necessary for RxL21 virulence function

To determine whether interaction with TPL is required for RxL21 virulence function, homozygous transgenic plants were generated expressing N-terminal HA tagged RxL21 or RxL21 $\Delta$ EAR in a Col-4 background under the control of the 35S promoter (HA:21#1/2 or HA:21 $\Delta$ EAR#1/2). Expression of the transgenes was verified by qPCR and western blot (S5 Fig). We noted that protein expression levels in all transgenic lines were very low perhaps due to a detrimental effect of effector accumulation. However, given the effector mRNA accumulation, we challenged these lines, compared to Col-4 WT and HA:GFP (35S promoter, Col-4 background) control plants, to *Hpa* infection. HA:21 lines exhibited enhanced susceptibility whereas HA:21 $\Delta$ EAR lines did not differ in susceptibility from WT and GFP controls (Fig 3A). Similarly, compared to HA:GFP controls, both HA:21 lines showed significantly enhanced lesion area at 72 hpi after inoculation with *B. cinerea* whereas HA:21 $\Delta$ EAR lines did not (Fig 3B). Enhanced sporulation was also clearly seen on leaves from the HA:21#1 line compared to Col-4 WT, HA:GFP controls and HA:21 $\Delta$ EAR#1 line (S6 Fig). These data demonstrate that the EAR motif of RxL21 (and hence most likely interaction with TPL/TPR1) is necessary for enhanced susceptibility to biotrophic and necrotrophic pathogens.

We also generated lines expressing RxL21 and RxL21 $\Delta$ EAR under an estradiol-inducible promoter and with an N-terminal Myc tag in Col-0 background (Est:21#2/6/9 and Est:21 $\Delta$ EAR#1/2/3). Expression of RxL21 after estradiol induction was verified by qPCR and protein accumulation (before and after induction) was assessed by western blot (S7 Fig). After estradiol induction, Est:21 lines showed enhanced susceptibility to *Hpa* (Fig 3C) and *B. cinerea* infection (Fig 3D). In contrast, Est:21 $\Delta$ EAR lines did not show enhanced susceptibility to either pathogen, confirming that the EAR motif is necessary for the virulence function of RxL21, and that RxL21 expression shortly before infection is sufficient to enhance host susceptibility.

#### TPL family members function in defence against multiple pathogens

It has previously been shown that TPL, TPR1 and TPR4 function in defence against *Pst*, with redundancy observed between TPL and TPR1 [20]. We screened *tpr1-tpl-tpr4* triple knockout plants using *Hpa* isolate Noks1. The *tpr1-tpl-tpr4* plants were found to be significantly more susceptible (Fig 4A) than both Col-0 WT and Col-0 GUS control plants. In addition, *tpr1-tpl-tpr4* plants showed significantly increased lesion area after infection with *B. cinerea* (Fig 4B)



**Fig 3. The EAR motif is necessary for RxL21 virulence function** *in planta.* Transgenic lines expressing RxL21 under control of a 35S promoter and N-terminal HA tag (HA:21#1/2 and HA:21 $\Delta$ EAR#1/2) were compared to Col-4 WT and HA:GFP (Col-4 background) controls during infection with (A) *Hpa* isolate Noks1 (sporangiophore counts per seedling at 4 dpi). Letters indicate significance using a Kruskal Wallis test, p < 0.05, n = 45 (B) *B. cinerea.* Lesion area was measured at 72 hpi (letters indicate significance determined by a one way ANOVA, p < 0.05, n = 24). Error bars display 95% CE. (C) Estradiol inducible lines expressing RxL21 and RxL21 $\Delta$ EAR fused to an N-terminal Myc tag (Est:21#2/9 and Est:21 $\Delta$ EAR#1/2) were infected with *Hpa* 18 hours after induction with 30  $\mu$ M  $\beta$ -estradiol or mock treatment (n = 100 per treatment). (D) The same lines with estradiol or mock treatment were subsequently infected with *B. cinerea.* Lesion area was measured at 72 hpi (n = 30). Letters in (C) and (D) indicate significant difference between treatments using a 2-way ANOVA and Bonferroni's multiple comparison test. P < 0.05. Whiskers show data range. Experiments were repeated with similar results.

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and visual symptoms of sporulation (Fig 4C). These data suggest that transcriptional regulation by TPL and family members is a crucial part of immune signalling against multiple pathogens with different lifestyles.



Fig 4. TPL family members function in immunity against a biotrophic and a necrotrophic pathogen. (A) tpl-tpr1-tpr4 plants showed enhanced susceptibility to Hpa isolate Noks1 measured by sporangiophore counts per seedling at 4 dpi compared to Col-0 WT and 35s::GUS controls. Letters indicate significance using a Kruskal Wallis test, n = 45, p < 0.05. (B) Enhanced susceptibility of tpl-tpr1-tpr4 to B. cinerea infection was also observed as determined by lesion area at 48 hpi (n = 24, letters indicate significance using a one-way ANOVA and a Tukey test, p < 0.05) and (C) visual inspection of sporulation. Here we show representative leaf images at 96h post infection. Scale bar is 1 cm.

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#### **RxL21 also interacts with TPR1**

TPL/TPR interactions can exhibit both redundancy (eg. all family members interact with multiple IAAs) and specificity (eg. several ERF TFs interact with 1 member or a subset of the TPL/ TPR family [23]). We tested interaction between all four TPRs and RxL21 using Y2H. In addition to TPL, RxL21 was found to interact with TPR1 (S8A Fig), the most closely related family member to TPL with 95% similarity at the amino acid level [20]. As with TPL, this interaction was abolished when the EAR motif of RxL21 was deleted. The RxL21-TPR1 interaction was also confirmed by Co-IP (S8B Fig) using transient expression in *N. benthamiana*.

#### RxL21 interaction with TPL affects host transcription

Both the nuclear localisation and protein-protein interaction with TPL within the nucleus suggest that RxL21 plays a role in manipulating host transcription. We observed that when GFP:: RxL21 is transiently expressed in *N. benthamiana*, treating with DAPI before imaging results in co-localisation of GFP and DAPI (S9 Fig). DAPI has been shown to accumulate with high levels of heterochromatin [54] and therefore tightly bound chromatin in a repressed state. We therefore speculated that RxL21 co-localisation with DAPI is indicative of an effector mechanism involving transcriptional repression via interaction with TPL. Consequently, we performed RNA-sequencing to compare gene expression in HA:21 and HA:21 $\Delta$ EAR lines, before and after induction of PTI with flg22; a conserved stretch of 22 amino acids from bacterial flagellin that is recognised by the receptor FLS2 [55]. We compared two independent HA:21 lines with two HA:21 $\Delta$ EAR lines, 2 h after both mock treatment and flg22 induction (full details of lines and treatments used for each sample is included in S2 Table). The total number of 75 bp paired-end reads generated was 676,059,072 across 24 samples. Transcript abundance was calculated by pseudoalignment of reads to AtRTD2 (Arabidopsis reference transcript dataset) [56] using Kallisto [57]. Genes with low expression across all samples were removed and TMM normalization was performed [58]. RNA count data after filtering and normalisation is provided in S12 Table.

A PCA plot was used to visualise the data quality and variance between conditions, using the average read counts of biological replicates (S10 Fig); samples from independent transgenic lines (HA:21#1/2 or HA:21 $\Delta$ EAR#1/2) were treated as biological replicates. 49% of variance between the samples can be explained by PC1 (flg22 treatment) however separation between RxL21 and RxL21 $\Delta$ EAR lines is also observed. Differentially expressed genes (DEGs) were defined as having a log<sub>2</sub> fold change of 1 or greater, with an adjusted p-value cut-off (after false discovery rate correction) of 0.05. DEGs are listed in <u>S3 Table</u>; the total number of DEGs for each comparison is shown in Fig 5A.

#### RxL21 does not perturb the overall PTI response

The scale of transcriptional reprogramming in response to flg22 treatment is similar in both the HA:21 and HA:21 $\Delta$ EAR lines, and indeed the vast majority of the flg22-induced DEGs are conserved across the two genotypes (2681 genes, with direction of differential expression also conserved: 591 downregulated and 2090 upregulated). Furthermore, DEGs induced upon flg22 treatment in HA:21 and HA:21 $\Delta$ EAR lines are strongly positively correlated with DEGs in Col-0 at 120 minutes post-flg22 treatment (compared to water treatment) from Rallapalli et al. (2014) (S4 Table; Pearson correlation coefficients of 0.841 and 0.844 respectively). Hence the majority of the flg22 response is intact in HA:21 lines and RxL21-induced pathogen susceptibility is not due to large-scale interference with the PTI response.

#### RxL21 causes differential expression of defence-related genes

There were only 417 genes differentially expressed (DE) between RxL21 and RxL21 $\Delta$ EAR transgenic lines under mock conditions, and 240 after flg22 treatment (Fig 5A; S3 Table). 113 genes were DE in both of these conditions, with the direction of differential expression conserved (Fig 5B). We analysed the DEGs in two groups: those apparent under mock conditions (417 genes, 316 up and 101 down) and the subset of DEGs specific to flg22 treatment (127; 68 up and 59 down) (Fig 5B). In both the genes downregulated and upregulated by RxL21 compared to RxL21 $\Delta$ EAR (under mock conditions) we found significant over-representation of Gene Ontology (GO) terms involved in response to biotic stimulus/defence and secondary metabolism. Enzymatic activity including oxidoreductase and hydrolase activity, and components of the endomembrane system were also enriched across the DEGs in both mock and flg22 treatments (S5 Table).

A significant proportion of the genes whose expression is perturbed by the RxL21 effector are associated with plant immunity (S6 Table). 82 genes differentially expressed in response to flg22 in WT plants [59] are mis-regulated by the effector, either before or after flg22 treatment. 213 genes differentially expressed in response to RxL21 are differentially expressed during *Pst* infection (DC3000 and/or DC3000HrpA compared to mock) [13], and 142 during *B. cinerea* infection [60].



Fig 5. RNAseq identifies differentially expressed genes in RxL21 compared to RxL21 $\Delta$ EAR transgenic lines. RNAseq was performed on HA-tagged RxL21 and RxL21 $\Delta$ EAR-expressing transgenic lines. (A) The number of upand down-regulated genes among differentially expressed genes under mock and flg22 treatment. DEGs were defined as having a log<sub>2</sub> fold change  $\geq 1$  or  $\leq$ -1, and a BH adjusted p-value of < 0.05. (B) Venn diagram shows differentially expressed genes between RxL21 and RxL21 $\Delta$ EAR after mock and flg22 treatment with an overlap of 84 upregulated and 29 down regulated genes. (C) Eight genes were selected for validation in estradiol (Est) inducible RxL21 and RxL21 $\Delta$ EAR expressing lines and Col-0 WT. We treated Arabidopsis seedlings with 30  $\mu$ M of  $\beta$ -estradiol for 18 hours. For Est:21 and Est:21 $\Delta$ EAR, data were obtained from 2 independent lines each with 3 biological replicates and expression was normalised to Arabidopsis tubulin gene. Black circles are individual data points and bars denote the mean  $\pm$  SE of target gene expression. Letters indicate significant differences (P < 0.05) (One-way ANOVA with Tukey's honest significance difference).

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We identified specific cases where RxL21 appeared to be suppressing the host defence response. Fifteen genes that are induced in response to flg22 treatment in WT Arabidopsis [59] were downregulated by the full-length effector compared to control after flg22 induction (including *AVRPPHB SUSCEPTIBLE 3*, *CRK24*, *CRK38* and *MYB85*). The *Pst* data set profiles gene expression in both virulent *Pst* DC3000 (capable of secreting effector proteins directly

into plant cells via the Type III secretion system) as well as a strain DC3000HrpA- which lacks the Type III secretion system. Hence, we can distinguish between host response and pathogen manipulation of gene expression via effector proteins. 20 defence genes that are suppressed by *Pst* effectors were regulated in the same manner by RxL21 (i.e. both *Pst* effectors and RxL21 downregulated or upregulated expression compared to respective controls). Furthermore, 18 genes that are specifically expressed in response to *Pst* effectors (and not part of the host PTI response) were also differentially expressed in response to RxL21. These data indicate not only that RxL21 DEGs are involved in plant immunity, but also that shared mechanisms of host manipulation exist between RxL21 from *Hpa* and *Pst* effectors.

We quantified expression of 8 DEGs using qPCR in an independent set of lines; estradiolinducible lines Est:21, Est:21 $\Delta$ EAR and Col-0 control. We selected eight of the genes downregulated by RxL21 compared to RxL21 $\Delta$ EAR; these genes code for two TFs (WRKY63 and NAC019), three receptor-like proteins, two calmodulins and a protein of unknown function (Sheet B in <u>S6 Table</u>). Seven out of the eight genes were significantly downregulated in Est:21 compared to Est:21 $\Delta$ EAR lines (Fig 5C). Six of these genes also showed reduced expression in Est:21 compared to wildtype Col-0. NAC019 did not differ between Col-0 and Est:21 but was reduced in Est:21 $\Delta$ EAR between Est:21 $\Delta$ EAR, and WRKY63 showed an intermediate expression level in Est:21 $\Delta$ EAR between Est:21 and Col-0. In general, the qPCR shows a similar pattern of log<sub>2</sub> fold change for the eight selected genes between RxL21 and RxL21 $\Delta$ EAR lines compared to RNA-seq results (S11 Fig).

#### RxL21-repressed genes are enriched for TPR1 binding targets

We hypothesised that RxL21 modulates host gene expression via interaction with TPL/TPR1. RxL21 could recruit these corepressors to new sites on the genome (by either binding to the DNA itself or to TFs), or RxL21 might sustain TPL/TPR1 repression when it would normally be relieved during infection. Using iDNA-Prot [61] we found no evidence to suggest that RxL21 contains a DNA binding motif. Promoters of DEGs under mock conditions were significantly enriched for WRKY TF binding motifs in both up and down-regulated genes (S7 Table). In the DEGs only evident after flg22 treatment, we found significant enrichment for MYB TF binding motifs (as well as a WRKY binding motif) in the upregulated genes and CAMTA (Calmodulin-binding transcription activator) binding motifs in the genes downregulated by RxL21 (S7 Table). This suggests that RxL21 may exert at least some of its effects via modulating TPL/TPR interaction with WRKY, MYB and CAMTA TFs.

We also compared DEGs from our RNAseq analysis to chromatin immunoprecipitation (ChIP)-seq data revealing TPR1 target/bound genes from *pTPR1:TPR1:GFP* Col plants (Griebel et al. [62], Fig 6A). Only genes repressed by RxL21 compared to RxL21ΔEAR (both under mock and flg22 treatment) were enriched for TPR1 targets (Sheet A, S8 Table) with binding observed immediately upstream of the transcription start site (TSS) (Fig 6A, purple and orange lines). Hence, for at least some of the RxL21 DEGs, RxL21 appears to impair expression by interaction with TPR1 upstream of the TSS. As expected, the level of TPR1 ChIP signal in RxL21-repressed DEGs was lower than in the group of all genes defined as TPR1 targets from the ChIP-seq data. The 20 genes repressed by RxL21 and identified as TPR1 targets encode several known positive regulators of immunity against biotrophic pathogens (*PBS3, ICS1* and *RLP20*) and regulators of abiotic stress tolerance (*WRKY46, WRKY63, NAC019*) (S8 Table). The identification of *WRKY46* and 63 as likely direct targets of RxL21 DEGs, suggests that these two TFs could be one key mechanism underlying RxL21-induced pathogen susceptibility. Another key mechanism appears to be mis-regulation of salicylic acid (SA) signalling with



Fig 6. Genes that are bound by TPR1 are overrepresented in genes repressed by RxL21. (A) Arabidopsis genes with repressed expression in HA:RxL21 lines compared to HA:21 $\Delta$ EAR lines show weak TPR1 binding. Metaplots display the TPR1 enrichment around the transcription start site (TSS) on genes regulated by RxL21 (BH adjusted p-value < 0.05 and log2-fold-change  $\geq$ 1, with or without flg22) or control genes without evidence for RxL21 dependent expression regulation (RxL21 control). TPR1 bound genes defined in Griebel et al. (2020) were used as a positive control (red line). On the y-axis is mean read count for the TRP1-GFP ChIP samples normalized to the input samples. TPR1 ChIP and input samples were scaled based on the number of mapped reads. TES = transcription end site. (B) ChIP-qPCR of RxL21-repressed genes. Two-week old seedlings, overexpressing RxL21 or RxL21 $\Delta$ EAR with an N-terminal myc tag and under the control of an estradiol inducible promoter, were treated with 30  $\mu$ M  $\beta$ -estradiol to induce *RxL21* expression 18 hrs before harvesting and cross-linking with 1% formaldehyde. ChIP assays were performed with an anti-Myc antibody. In the ChIP-qPCR, the enrichment of immunoprecipitated DNA was normalized by the percent input method (signals obtained from ChIP samples were divided by signals obtained from an input sample). Error bar represents ± SE of four technical repeats. Arabidopsis *Actin 2* was included as a control but no amplification was detected after 40 cycles. The experiment was repeated with similar results.

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RxL21/TPR1 targets including genes for two enzymes required for SA accumulation (ICS1 and PBS3).

Genes that do not respond to the effector (RxL21 control in Fig 6A; Sheet B in S8 Table) do not show enrichment for TPR1 binding, while interestingly, genes induced by RxL21 (compared to RxL21 $\Delta$ EAR in mock or after flg22 treatment; Fig 6A, yellow and brown lines) demonstrate depletion of TPR1 binding around the TSS. This suggests that promoters of genes upregulated by RxL21 are not TPR1 targets and mis-regulation of these by the effector is likely to be an indirect effect rather than through mis-placing bound TPR1.

To confirm the observed overlap of RxL21 repressed genes with TPR1 binding sites, we performed independent ChIP-qPCR in Est:21 and Est:21∆EAR lines (Fig 6B) on seven genes (marked in bold in <u>S8 Table</u>). Promoter regions encompassing 500 bp upstream and 500 bp down-stream of the TSS for five of these genes (*NAC019*, *AED1*, *STMP6*, *AT5G44574* and *HR4*) were enriched approximately 5-fold in RxL21 immunoprecipitated samples compared to RxL21∆EAR samples. The promoter regions of *CRK38* and *WRKY63* didn't show strong enrichment, however we noted that *CRK38* was only repressed by RxL21 after flg22 treatment in our data set. We also amplified *AtActin2* as a negative control which was not detectable in ChIP samples after 40 qPCR cycles. Promoter regions of these five genes can, therefore, be directly bound by both RxL21 (and not RxL21 $\Delta$ EAR) and TPR1. Furthermore, a lack of enrichment in the ChIP-qPCR assays in RxL21 $\Delta$ EAR samples suggests that RxL21 binds to these promoter regions via TPR1 (maintaining repression when it would normally be relieved) rather than recruiting TPR1 to new locations.

# Discussion

#### RxL21 mimics a host gene regulation mechanism

Manipulation of host transcription by effector proteins is a key mechanism by which plant pathogens are able to render the plant a more hospitable environment for colonization. In this study we provide evidence that an effector from the oomycete pathogen Hpa mimics a mechanism by which plants regulate transcription throughout their development and in response to stress; the EAR motif is a repression motif that occurs in plant transcriptional repressors and mediates interaction with members of the TPL/TPR corepressor family. This mechanism of TPL recruitment to transcriptional repression complexes is highly conserved across the plant kingdom [26,34,36]. The Hpa effector RxL21 contains a C-terminal EAR motif which we have shown to be responsible for interaction with the CTLH domain of the transcriptional corepressor TPL in both *in vitro* yeast assays and *in planta* (Co-IP and Split YFP; Figs 2 and S3). This interaction is specific, and is abolished even when Leu residues in the EAR motif are substituted with the similar amino acid Ile (Fig 2 and S2 Fig). A pivotal role for TPL family members in regulation of immune responses has been previously implicated [20,63] and Arabidopsis plants lacking expression of TPL and its closest homologues, TPR1 and TPR4, show enhanced susceptibility to Pst, Hpa and B. cinerea (Zhu et al., 2010 and Fig 4). Interestingly, the effector RxL21 also interacts with TPR1 (S8 Fig) and, when expressed in planta, enhances susceptibility of Arabidopsis to a biotrophic (Hpa), hemibiotrophic (Pst) and necrotrophic pathogen (B. cinerea)(Figs 1, 3 and S6). There are a few examples of EAR motifs in other pathogen effectors -the bacterial XopD effector family [41] and an effector from fungi that contains an EAR motif and interacts with TPR4 [44]. Crucially we show that the C-terminal EAR motif (and thus interaction with TPL/TPR1) is required for enhanced pathogen susceptibility provided by the RxL21 effector (Fig 3).

#### RxL21 alters expression of a subset of the host defence response

By performing RNA-seq comparing the transcriptional effects of expressing RxL21 compared to the effector lacking the EAR motif (RxL21 $\Delta$ EAR) we were able to identify specific DEGs that result from RxL21-TPL/TPR1 interaction (Fig 5). Importantly, there is no large-scale change in gene expression in response to presence of the effector (either with or without flg22 treatment to induce PTI), yet effector expression still delivers a striking susceptibility enhancement to the host plant. This is similar to the situation with *Pst* where blocking of effector delivery through the Type III Secretion System only perturbed the expression of 872 PTI-regulated genes with the vast majority (5350) showing no change in expression or amplification of usual PTI-regulation [13]. These findings highlight the ability of pathogen effectors to target (and hence identify) specific components of the plant defence response with great effect.

The genes mis-regulated by RxL21 include several important components of plant defence responses. These include down regulation of several receptor-like protein (RLP)genes; *RLP6*, *RLP22*, *RLP23* and *RLP35*, with *RLP20* DE specifically after flg22 treatment. RLPs are the second largest family of Arabidopsis leucine rich repeat-containing receptors; several members

have been shown to function as PAMP receptors and it is becoming increasingly clear that RLPs play a critical role in pathogen recognition [64]. Consistent with this, many *RLP* genes (including *RLP6*, *22*, *23* and *35*) show increased expression during pathogen infection and/or hormone treatment [64,65]. Several RLPs are involved in fungal and oomycete resistance [66–70]. In addition, several other genes downregulated by RxL21 are known components of the defence response. These include genes that impact resistance against biotrophic pathogens such as *accelerated cell death* 6 (*ACD6*) and Cysteine-rich receptor-like kinase *CRK13* [71] that positively regulate SA signalling [72], as well as Calmodulin-like *CML41* that regulates flg22-induced stomatal closure [73]. Furthermore, miR825 target *AT3G04220* [74] and the lipid transfer protein AZI3 [75] contribute to defence against necrotrophs and the lectin receptor kinase LecRK4.1 positively regulates Arabidopsis PTI and resistance against biotrophic and necrotrophic pathogens [76]. Reduction in expression of these genes is likely to contribute to the observed enhanced susceptibility of RxL21 over-expressing plants to both biotrophic and necrotrophic pathogens.

Twenty genes that are direct TPR1 targets are repressed by RxL21, seven of which were specific to flg22 treatment. This includes tetraspanin9 (*TET9*) which was repressed 2.6-fold by RxL21 compared to RxL21 $\Delta$ EAR. TET8/9 promote the formation of exosome-like extracellular vesicles to deliver host sRNAs into *B. cinerea* to decrease fungal virulence [77]. TET9 accumulates around fungal infection sites after *B. cinerea* infection and *tet9* loss-of-function mutants display weak but consistent enhanced susceptibility towards *B. cinerea* [77], leading us to speculate that *TET9* repression could contribute to the *B. cinerea* susceptibility we see in RxL21-expressing lines.

Two WRKY TFs (WRKY46 and WRKY63) are direct TPR1 targets and repressed in RxL21 expressing plants in comparison to the RxL21 $\Delta$ EAR variant. WRKY TFs have been shown to regulate defence responses against biotrophic and necrotrophic pathogens [78,79] and WRKY46 overexpressing plants are more resistant to *Pst* [80]. Moreover, non-host resistance against *Erwinia amylovora* is regulated by WRKY46 and WRKY54 via EDS1 [81] and WRKY46 is a transcriptional activator of *PBS3* expression [82]. Consistent with this, *PBS3* expression is also reduced in RxL21 compared to RxL21 $\Delta$ EAR over-expressing plants. Recently, it was shown that PBS3 binds to and protects EDS1 from proteasomal digestion [83], hence reductions in PBS3 expression could lead to lower levels of EDS1 protein and enhanced susceptibility to biotrophic pathogens [84]. Both WRKY46 and 63 bind to the promoter of the SA biosynthesis gene *SID2* with increased levels of SA, correlated with increased levels of expression of these and several other WRKY TFs [85]. Consistent with this, *SID2* (also called *ICS1*) is downregulated by RxL21 after flg22 treatment, however this may also be due to direct repression via association with TPR1 as *SID2* is also a target of TPR1.

WRKY TF binding motifs were overrepresented in the promoters of RxL21 DEGs implying that many of the DEGs are downstream targets of WRKY TFs. It is possible that WRKY46 and WRKY63 are directly repressed by RxL21 which in turn leads to changes in expression of their target genes. We observed overrepresentation of CAMTA motifs in genes that were specifically downregulated by RxL21 in an EAR-dependent manner upon flg22 treatment, indicating that target genes of CAMTA TFs are showing differential expression. There is no evidence to date that CAMTA TFs are TPL/TPR targets (Causier, Ashworth, Guo, & Davies, 2012a) but CAMTA TFs play a role in immune regulation through suppressing pathogen-responsive genes and could therefore be important targets for RxL21 manipulation via TPL/TPR1 [86–88].

#### Recruitment of RxL21 to TPL/TPR1 transcriptional complexes

We have shown that the suppression of immunity by RxL21 depends on its EAR domain, and hence most likely through modifying actions of TPL and TPR1. Several scenarios are possible.

On the one hand, RxL21 could interact with a TF and mediate subsequent TPL/TPR1 binding (to novel sites) in a manner similar to NINJA or JAZ proteins [25]. Evidence for an interaction of RxL21 with the TF TCP14 was seen in Y2H experiments [17] and later it was shown that TCP14 shifts RxL21 into subnuclear foci [18]. However, if RxL21 was binding TFs independently of TPL/TPR1 then our ChIP-PCR experiments would be expected to show a similar enrichment in the RxL21 and RxL21AEAR at the tested loci. An alternative model is that RxL21 interferes with the repression (or lifting of repression) of existing TPL/TPR1 targets. We provide evidence to show that, at least for some genes, RxL21 appears to bind to TPL/ TPR1 within transcriptional complexes at plant gene promoters and prevent transcriptional de-repression (Fig 6). First, there is a significant over-representation in TPR1 binding sites upstream of genes that are repressed by full-length RxL21 compared to RxL21ΔEAR (where interaction with TPL/TPR1 is lost by deletion of the EAR motif). Second, promoters of several RxL21-repressed genes were shown to be binding targets of TPR1 and RxL21 (and not of  $RxL21\Delta EAR$ ). This indicates that in at least these cases, RxL21 likely binds to TPR1 protein already present at the gene promoter (as there is no binding of RxL21 to these promoters without the EAR motif) and subsequently exerts its activity on the TPL/TPR1 complex.

The TPL family of proteins are closely related to Groucho / Tup1 proteins in animals and fungi [89] which, like TPL, have been shown to form tetrameric structures. Tetramerization was suggested as a mechanism to enable recruitment of multiple TFs to a single complex [30] and binding of EAR motif peptides does not prevent tetramerization [30,90,91]. It is therefore plausible that RxL21 is able to bind TPL/TPR1 via its EAR motif while other epitopes of the TPL oligomer are binding other proteins such as TFs within the transcriptional complex. How TPL/TPR1-bound RxL21 behaves is not known. However, TPR1 activity was shown to be regulated by the (SUMO) E3 ligase SIZ1 [92], perhaps via SUMOylation inhibiting the corepressor activity of TPR1 by preventing its interaction with HISTONE DEACETYLASE 19 (HDA19). It is possible that RxL21 is shielding the SUMO attachment sites K282 and K721 in TPR1 and/or preventing (SUMO) E3 ligase activity, thereby enhancing the TPL/TPR corepressor activity.

There is no evidence that the only mechanism of RxL21 action is through maintained repression of direct TPL/TPR1 targets and it is worth noting that many DEGs were upregulated in the presence of RxL21. In addition to immune suppression, pathogen effectors are known to directly manipulate plant gene expression, and hence physiology, to aid infection [93,94]. During *Pst* infection, more than 1500 genes were specifically upregulated in response to *Pst* effectors, and we observed an overlap (15 genes) between these genes and those upregulated by RxL21. However, it is possible that upregulated genes are downstream targets of TFs or other regulators targeted by RxL21. This appears to be the most likely explanation given that the RxL21 upregulated genes are not enriched for TPR1 binding in wildtype plants and the enrichment for WRKY TF binding motifs in the promoter regions of these genes.

Remarkably, *in planta* expression of RxL21 alone can increase susceptibility to pathogens with varying lifestyle and virulence strategies. To our knowledge there are very few, if any, effectors that exhibit this activity. RxL21 is one of several effectors that mimic the EAR motif for transcriptional repression, and appears to actively initiate and/or maintain repression of gene expression mediated by TPL/TPR1. We show here that the RxL21 EAR motif is essential for its virulence function, and for modifying the expression of key host defence genes. Future interrogations will be to examine the RxL21 mode of action on TPL/TPR1 transcriptional repression complexes and determine which effector-manipulated defences ultimately result in enhanced susceptibility of the host plant.

## Methods

#### Sequence alignment

Alignment of HaRxL21 alleles was performed using sequences in Asai et al. [45] or BioProject PRJNA298674 (for Noks1). Alignment was performed using T-COFFEE [95] (Version\_11.00. d625267; http://tcoffee.crg.cat/apps/tcoffee/do:mcoffee). SignalP-5.0 (http://www.cbs.dtu.dk/ services/SignalP/) was used to predict the signal peptide cleavage site.

# Yeast-2-Hybrid

Yeast-2-Hybrid (Y2H) screening was performed as described in Dreze et al. [96]. Briefly, yeast strains Y8800 and Y8930 harbouring AD-X and DB-X constructs respectively were mated on yeast-extract, peptone, dextrose, Adenine (YPDA) medium. Yeast was replica plated onto Synthetic complete (SC) media lacking combinations of Leu, Trp, His and Ade. 3AT (3-amino-1,2,4- triazole) was added to selective plates lacking His to increase stringency as a competitive inhibitor of the HIS3 gene product. Plates were cleaned using sterile velvets after 1 day and imaged after 4 days. To determine that constructs were not autoactivating, AD-X and BD-X constructs were mated with Y9830 and Y8800 harbouring empty pDB and pAD respectively and any autoactivating constructs were removed from further screening. To generate EAR motif mutants, site-directed mutagenesis (SDM) was performed using a QuikChange II Site-Directed Mutagenesis Kit (Agilent, Santa Clara, US). Primer sequences for SDM are included in S9 Table.

# **Plant material**

35S::HaRxL21 lines (RxL21a/b) are described in Fabro et al. [16]. To generate 35S::HA:: HaRxL21 lines in Arabidopsis, RxL21 and RxL21ΔEAR were cloned into pEarleygate201 [97] and transformed into Arabidopsis ecotype Col-4 using floral dipping {Clough:1998vw}. Independent transformants were selected on BASTA (Bayer CropScience, Wolfenbüttel, Germany) until homozygous. Western Blotting was performed on 14 day old seedlings to determine protein expression using anti-HA high affinity antibody (Roche, Penzberg, Germany). Primer sequences for RxL21 and RxL21 deletion variants are in S9 Table. To generate estradiol inducible lines, RxL21 and RxL21ΔEAR were cloned into the pER8 plasmid {Zuo:2000} and transformed into Arabidopsis (Col-0 background) via floral dip. Independent homozygous lines were selected on hygromycin (Invitrogen, Carlsbad, US).

# **Pathogen Assays**

*Hpa* infection screens were performed as described in Tomé et al. [98]. Briefly, plants were sown in a randomized block design within the inner modules of p40 seed trays at a density of approximately 30 seedlings per module. Modules at the periphery of the tray were sown with WT. Plants were grown under a 12 h photoperiod, 20°C, 60% humidity. *Hpa* isolates Noks1 and Maks9 were maintained on Col-0 plants and inoculated onto 2 week old seedlings at  $3x10^4$  spores/mL. After infection, plants were covered with a lid and sealed. Sporangiophore counts were performed using a dissecting microscope at 4 dpi. Botrytis infection screens were performed as described in Windram et al. [60]. Briefly, *B. cinerea* (strain pepper) was maintained on sterile tinned apricot halves in petri dishes, kept in the dark at 25°C. Detached Arabidopsis leaves were placed on 0.8% agar and inoculated with a 10 μL droplet of spore suspension at  $1x10^5$  spores/mL in 50% sterile grape juice. Lesions were imaged at 24, 48, 72 and 96 h post infection (hpi). For Botrytis infections on estradiol-inducible lines, plants were grown at 10 h photoperiod, 16°C at 60% humidity. Gene expression was induced by 30 μM β-

estradiol, 18 h before infection. Spores were washed from the surface of the *B. cinerea* culture plate using potato dextrose broth (PDB) medium and leaves were infected using a 5  $\mu$ L droplet at a concentration of 1x10<sup>4</sup> spores/mL.

#### Localisation and Split YFP

For localisation, pK7WGF2 (N-terminal GFP fusion; [99]) and pB7WGR2 (N-terminal RFP fusion; [100]) vectors were used. BiFP (Bimolecular Fluorescence complementation in Planta) vectors were used for generation of C- or N- YFP fusion constructs [101]. Expression vectors were transformed into *A. tumefaciens* strain GV3101 and cultured overnight at 28°C. *A. tumefaciens* harbouring expression constructs, along with P19 [102] were co-infiltrated into *N. benthamiana*. Leaves were imaged 48 h after infiltration by laser scanning confocal microscopy.

#### **Co-immunoprecipitation**

For Co-immunoprecipitation (Co-IP), TPL and RxL21 were expressed in pCsVMV-HA3-N-1300 (C-terminal HA tag) and Per8 (N-terminal myc tag) vectors. Proteins were expressed transiently in N. benthamiana. Leaves of about four-week-old plants were infiltrated with A. tumefaciens (OD600 = 0.2) in infiltration buffer (10 mM MgCl<sub>2</sub>, 10 mM MES, and 100  $\mu$ M acetosyringone). After 24 hpi, leaves were sprayed with 30  $\mu$ M estradiol to induce the expression of RxL21 and subsequent mutants. At 48 hpi, approximately 2 g of tissue from the infiltrated area was collected and frozen with liquid N<sub>2</sub>, then ground into powder using a mortar and pestle. All of the steps were carried out either on ice or in a 4°C cold room. About three volumes of nuclei extraction buffer (NEB) (20 mM Hepes pH 8, 250mM Sucrose, 1mM Magnesium Chloride, 5mM Potassium Chloride, 40% (v/v) glycerol, 0.25% Triton X-100, 0.1 mM PMSF, Protease inhibitor cocktail (Roche cOmplete), were added to each sample. The resuspended samples were filtered using miracloth and centrifuged at 3,320 g for 15 min at 4°C. The pellet was subsequently resuspended and washed with NEB in 2 mL microcentrifuge tubes. Washed pellet was resuspended in lysis buffer (10 mM Tris pH7.5, 0.5 mM EDTA, 1 mM PMSF, 1% Protease Inhibitor, 150 mM Sodium Chloride, 0.5% Igepal) and sonicated to break the nuclei. After sonication, the supernatant was centrifuged again to remove additional debris, and the supernatant was used as input for IP. Afterwards, the µMACS kit of magnetic microbeads, conjugated to an anti-c-myc monoclonal antibody (130-091-123; Miltenyi Biotech), was used for IP according to the manufacturer's instructions. Eluted proteins were analysed by western blot using HRP-conjugated anti-myc antibody (130-092-113; Miltenyi Biotec), or an anti-HA antibody (3F10; Roche).

## **RNA-Seq**

**Sample preparation.** Arabidopsis seedlings were grown on ½ MS agar with sucrose for 14 days at 22 °C with photoperiod of 8 h light and 16 h dark, and 150 mmol photons \*  $m^{-2} \times s^{-1}$ . Seedlings were then transferred to 6 well plates containing 5 mL ½ MS liquid media (approximately 8 seedlings per well) and left to rest overnight. Media was replaced with 5 mL fresh ½ MS liquid media containing 100 nM flg22 or 5 mL ½ MS control. Each sample consisted of total pooled seedlings from a well, these were harvested 2 h post induction, briefly dried on tissue and flash frozen in liquid nitrogen. For each of 35S::HA::RxL21 and 35S::HA::RxL21\DeltaEAR, two independently transformed Arabidopsis lines were used (HA:21#1/2 and HA:21\DeltaEAR#1/2), each with 3 biological replicates. RNA was extracted from Arabidopsis tissue using Trizol (Invitrogen) and cleaned up using a RNeasy Mini Kit (Qiagen), Libraries were made

using the NEBNext Ultra Directional RNA library prep kit. Each library was sequenced on two lanes of Illumina HiSeq 4000 generating 75 bp paired end reads. The sequencing files were examined using fastQC version 0.11.5.

Quantification of transcripts and DE gene analysis. Transcript abundances were generated using Kallisto version 0.44.0 [57] using an index generated using Arabidopsis Thaliana Reference Transcript Dataset 2 [56]. The number of bootstraps was set to 100. DEGs were generated using the 3D-RNA-seq app [103]. Sequencing replicates from the two HiSeq 4000 lanes were merged. Data was pre-processed by filtering to remove genes which did not meet the following criteria; 1) An expressed transcript must have at least 3 out of 23 samples with CPM (count per million read)  $\geq$ 1 and 2) An expressed gene must have at least one expressed transcript. One sample was removed from further analysis ( $RxL21\Delta EAR.flg22$  line 2, biorep 3) due to outlying sequencing depth. RNA-seq read counts were normalised with Trimmed Mean of M-values method [58]. Models of expression contrasts (RxL21.flg22 vs RxL21.mock, RxL21\DEAR.flg22 vs RxL21\DEAR.mock, RxL21.mock vs RxL21\DEAR.mock and RxL21.flg22 vs RxL21 $\Delta$ EAR.flg22) were fitted using Limma Voom [104,105]. Genes were significantly DE in the contrast groups if they had BH adjusted p-value < 0.05 and  $\log_2$ -fold-change > 1. We performed Go-Term analysis for biological process, cellular component and molecular function using AgriGO ([106], http://bioinfo.cau.edu.cn/agriGO/). Significance of GO enrichment was determined with FDR < 0.05. Known Arabidopsis TF DNA-binding motifs were retrieved from CIS-DB version 1.02 [107], and those described in Franco-Zorilla et al., [108]. Promoter sequences defined as the 500 bp upstream of the transcription start site were retrieved from https://www.arabidopsis.org/ (Araport 11 annotation). Motif occurrences were determined using FIMO [109], and promoters defined as positive for a motif if it had at least one match with a P value  $< 10^{-4}$ . Motif enrichment was assessed using the hypergeometric distribution against the background of all genes. *P* values < 0.001 were considered significant to allow for multiple testing. RNA seq data is available online at NCBI under Bioproject ID: PRJNA622757.

# qPCR

Total RNA extraction was performed with TRIzol (Invitrogen) following the manufacturer's instructions. cDNA was synthesized by using the RevertAid cDNA synthesis kit (Thermofisher Scientific, Waltham, US). Quantitative PCR was performed in the Applied Biosystems 7500 FAST real-time PCR system using SYBR Green JumpStart Taq ReadyMix (Sigma-Aldrich, St Louis, US). Transcript levels of target genes were determined via the 2- $\Delta\Delta$ Ct method [110], normalized to the amount of *Arabidopsis tubulin 4* (*AT5G44340*) transcript. For expression analysis of HA:RxL21 and HA: 21 $\Delta$ EAR lines, expression was normalised to *Actin2* (*AT3G18780*) and *UBQ5* (AT3G62250) transcript levels. Primer sequences are detailed in <u>S9</u> Table.

#### Chromatin immunoprecipitation (ChIP) PCR

ChIP PCR was performed as described in Gendrel et al. [111]. Briefly, 2 week old seedlings were harvested and cross-linked in 1% formaldehyde (Sigma-Aldrich) solution under vacuum for 15 min. The isolated chromatin complex was resuspended in Nuclei lysis buffer 50 mM Tris-HCl, 10m M EDTA, % SDS and one tablet PI (Roche cOmplete) and sheared by sonication (Bioruptor, Diagenode, Ougrée, Belgium) to reduce the average DNA fragment size to around 500 bp. The sonicated chromatin complex was diluted in ChIP dilution buffer (16.7 mM Tris-HCl, 1.2 mM EDTA, 167 mM NaCl, 1.1% Triton X-100 and one tablet PI) and immunoprecipitated by anti-myc antibody (ab9132; ChIP-grade, Abcam, Cambridge, UK).

After reverse cross-linking, the immunoprecipitated DNA was extracted by using equal amounts of phenol/chloroform/isoamyl alcohol and precipitated with 100% EtOH, 1/10 volume of 3 M Sodium acetate and, 10 mg/mL glycogen. DNA was resuspended in Milli-Q water and analyzed by qPCR with gene specific primers. Primer sequences are shown in <u>S9 Table</u>. Input % in IP samples was calculated by (100\*2^(Adjusted input-IP)). *AtActin2 (AT3G18780)* was included as a control.

## Metaplots for TPR1 binding

Preprocessing of chromatin immunoprecipitation-sequencing (ChIP-seq) data for the *pTPR1*: *TPR1-GFP* Col-0 line [20] was performed as in Griebel et al. [62]. Briefly, adapters and other overrepresented sequences detected with fastqc (version 0.11.9; http://www.bioinformatics. babraham.ac.uk/projects/fastqc/) were removed with cutadapt (version 1.9.1, -e 0.2 -n 2 -m 30; [112]). Raw reads were mapped to *Arabidopsis thaliana* genome version TAIR10 (arabidopsis. org) with bowtie2 (version 2.2.8; [113]). The alignment files were filtered for low-quality reads (samtools view -q 10, version 1.9, [114,115]), deduplicated and merged from the three biological replicates. The metaplots were prepared with deepTools 3.3.0 following the manual pages (https://github.com/deeptools/deepTools/tree/develop). The TPR1-GFP ChIP-seq data were input-normalized using the bamCompare function (—operation subtract, default 'readCount' scaling).

# Supporting information

S1 Data. Excel spreadsheet containing, in separate sheets, the underlying numerical data and statistical analysis for Figures/Figure panels 1, 3, 4, 5c, 6b, S5a, S7a and S11. (XLSX)

S1 Fig. Amino acid sequence alignment of HaRxL21 alleles from *Hpa* isolates Cala2, Emco5, Emoy2, Emwa1, Hind2, Maks9, Noks1 and Waco9. Sequences were obtained from Asai et al., 2018 (Cala2, Emco5, Emoy2, Emwa1, Hind2, Maks9 and Waco9) and BioProject PRJNA298674 (Noks1). Multiple sequence alignment was performed using T-coffee (http:// tcoffee.crg.cat/apps/tcoffee/do:mcoffee). Sites of amino acid substitution between alleles are highlighted in white. Predicted signal peptide is shown in grey. The RxLR-DEER motif (green) and EAR motif (magenta) are conserved across alleles except Noks1. (TIF)

**S2 Fig. Site directed mutagenesis of Leucine residues in the EAR motif of RxL21 abolishes interaction with TPL by Yeast-2-Hybrid.** Leu residues in the EAR (LxLxL) motif were mutated to Phenylalanine (F), Alanine (A) and Isoleucine (I). Amino acids that differ from WT are indicated in Red. AD; GAL4 activation domain. DB; GAL4 DNA binding domain. (A) Interaction was tested against TPL from Arabidopsis (AtTPL). (B) Interaction was tested with TPL from *N. benthamiana* (NbTPL). Growth on media lacking Leu, Trp and His is shown, indicating successful mating and activation of the *GAL1::HIS3* reporter gene due to interaction. All combinations tested showed growth on media lacking only Leu and Trp indicating successful mating (see S1 Data). The experiment was repeated on multiple plates with similar results.

(TIF)

**S3 Fig. RxL21 interacts with TPL** *in planta* by BiFC. The BiFC assay was conducted transiently in *N. benthamiana* epidermal cells by co-agroinfiltration of  $TPL(YFP^N)$  and RxL21 ( $YFP^C$ ),  $TPL(YFP^N)$  and  $RxL21\Delta EAR(YFP^C)$ ,  $RxL21(YFP^N)$  and  $RxL21(YFP^C)$ ,  $TPL(YFP^N)$  and  $TPL(YFP^C)$ . Infiltrated tissues were imaged at 48 hpi by confocal scanning laser

microscopy for YFP fluorescence. The interaction between TPL and RxL21 was lost with deletion of the EAR motif. RxL21 does not appear to form dimers unlike TPL where strong fluorescence was observed in the nucleus. Scale bar =  $10 \mu m$ . (TIF)

**S4 Fig. Interaction with TPL is specific to RxL21 amongst** *Hpa* **EAR-motif containing effectors.** Y2H was performed using activation domain (AD)-TPL and DNA-binding domain (DB)-Effector constructs (top) and DB-TPL with AD-Effector constructs (bottom row). TPL interaction with HaRxL21 and TPL dimerisation were used as positive controls. Protein-protein interaction is shown by growth (indicating *GAL1::HIS3* reporter gene activation) on SC media lacking Leucine, Tryptophan and Histidine. All combinations tested also showed growth on–LW media indicating successful mating (see <u>S1 Data</u>). The experiment was repeated on multiple plates with similar results. (TIF)

**S5 Fig. Gene and protein expression in 35S::HA::RxL21 lines.** (A) Expression of *RxL21* by quantitative RT-PCR relative to *AtAct2* and *UBQ5*. Error bars show variance between technical replicates. Previously characterized 35S lines RxL21a/b were included for comparison. (B) Upper: Coomassie staining, lower: western blot using anti-HA. GFP and RxL21 bands are indicated by blue and red arrows respectively. (TIF)

**S6 Fig. RxL21 lines show more visual sporulation after** *Botrytis cinerea* infection. Photos taken 96 h post-infection with *Botrytis cinerea*. HA:21#1 appears to show more sporulation compared to Col-4, 35S::HA::GFP (HA:GFP) and HA:21 $\Delta$ EAR#1. Scale bar is 1 cm. (TIF)

S7 Fig. RxL21 expression in RxL21 and RxL21 $\Delta$ EAR estradiol inducible lines. (A) Relative expression of *RxL21* in Arabidopsis plants expressing myc::RxL21 and myc::RxL21 $\Delta$ EAR (under an estradiol (Est) inducible promoter) was determined by quantitative RT-PCR 18 h after induction with 30 µM estradiol. Expression levels were normalised to Arabidopsis *tubulin* <u>4</u>. Error bars show standard error between 3 biological replicates. (B) Anti Myc immunoblot showing Est-inducible expression of RxL21 and RxL21 $\Delta$ EAR in Arabidopsis lines. Samples were taken 18 h after induction with 30 µM estradiol. (TIF)

**S8 Fig. RxL21 interacts with TPR1.** (A) RxL21 interacts with TPL and TPR1 by Y2H, indicated by growth on media lacking Leucine (Leu), Tryptophan (Trp) and Histidine (His). Growth on–Leu-Trp media indicates successful mating. Y2H was repeated in both directions, with both RxL21 and TPRs fused to AD; activation domain and DB; DNA binding domain. Y2H was repeated on multiple plates with similar results. (B) TPR1 interacts with RxL21 *in planta*. TPR1:HA with Myc:RxL21 or Myc:RxL21 $\Delta$ EAR were transiently expressed in *N*. *benthamiana* leaves and harvested after 48 h. RxL21 expression was induced by 30 $\mu$ M  $\beta$ -estradiol 24 h prior to harvesting.  $\alpha$ -myc beads were used for immunoprecipitations (IP). HA antibody was used to detect TPR1 immunoblots (IB) and  $\alpha$ -myc antibody was used to detect RxL21 and RxL21 $\Delta$ EAR.

(TIF)

**S9 Fig. RxL21 co-localizes with DAPI in the nucleus.** Transient expression of GFP:RxL21 using Agrobacterium mediated transformation of *N. benthamiana*. Immediately prior to imaging, leaves were stained by infiltration with DAPI (4',6-diamidino-2-phenylindole). Scale bars

are 5 μm. (TIF)

**S10 Fig. Principal component analysis showing variance between samples used for RNA-seq.** A principal component analysis (PCA) plot characterizes the trends shown by the RNAseq data. Each point represents a sample (mean of 2 technical replicates) and colour characterizes the sample group. Groups consist of 6 bioreps (3 samples from each of two independent transgenic Arabidopsis lines expressing 35S:HA:RxL21 or 35S:HA:RxL21ΔEAR). (TIF)

S11 Fig. Comparison of RNA-seq expression data and qPCR data. Comparison of  $log_2$  fold change ( $log_2FC$ ) for the 8 selected genes between RxL21 and RxL21 $\Delta$ EAR lines by qPCR compared to RNA-seq read counts. Mean fold change of three biological replicates is shown. (TIF)

**S1 Table.** *Hpa* **RxLs and RxLLs found to contain the LxLxL motif.** Putative RxL and RxLlike (RxLL) effectors from *Hpa* were searched for the presence of a LxLxL motif. Effectors were characterised as having a C- or N-terminal EAR motif if the motif was detected within 35 amino acids of the N-or C-terminus of the effector sequence (otherwise 'mid'). Nuclear or cytoplasmic localisation of effectors as identified by Caillaud et al., (2011) is indicated. Effectors shown in <u>S4 Fig</u> and tested for interaction with TPL are highlighted in yellow. (XLSX)

**S2 Table. RNA Seq sample information and read counts.** (a) Line and treatment information for each sample. (b) Normalised read counts per gene after combining sequencing replicates, removing lowly expressed genes across all samples and normalisation using the limma-voom pipeline.

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(XLSX)
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**S3 Table. Differentially expressed genes after RNA-seq analysis.** Differentially expressed genes after RNA-seq analysis. Contrast indicates in which pair-wise comparison the gene is differentially expressed. Adj.pval indicates significance after limma-voom pipeline and Benjamini Hochberg false discovery rate correction. FC = log2 fold change. Arabidopsis gene descriptions are based on Araport 11 annotation. (XLSX)

S4 Table. Comparison of flg22 induced gene expression in RxL21 lines compared to Col-0 WT.  $Log_2$  fold expression of differentially expressed genes in RxL21 and RxL21 $\Delta$ EAR lines after flg22 treatment compared to mock, compared to  $Log_2$  fold expression in Col-0 at 120 minutes post treatment with flg22 compared to mock treatment from Rallapalli et al. (2014). (XLSX)

S5 Table. Over-represented GO categories in differentially expressed genes between RxL21 and RxL21 $\Delta$ EAR in both mock and flg22 conditions. Singular enrichment analysis of GO terms was performed using AgriGO. The cutoff P-value after false discovery rate correction is <0.05. (F = molecular function, P = biological process, C = cellular component). (XLSX)

**S6 Table.** (A) Genes differentially regulated in RxL21 compared to RxL21 $\Delta$ EAR. Gene descriptions are from Araport 11. Adj.pval indicates significance after limma-voom pipeline and Benjamini Hochberg false discovery rate correction. Log<sub>2</sub> fold change (FC) indicates expression in HA:RxL21 lines compared to HA:RxL21 $\Delta$ EAR lines. Flg22 specific DEGs only show significant differential expression after flg22 induction. Mock / flg22 independent DEGs

are DE under mock and/or mock and flg22 conditions. Flg22 (Column H) indicates expression values 120 minutes after flg22 induction from Rallapalli et al. (2014). Comparison is shown to *B. cinerea* responsive DEGs from Windram et al. (2012) and DEGs during *Pst* infection (Lewis et al. 2015), including (Column K) description of *Pst* expression type from Fig 4 (Lewis et al, 2015) where applicable. (B) Details of the genes used for qPCR for RNAseq verification. (XLSX)

S7 Table. Motifs Over-represented in RxL21 differentially expressed genes identified using RNA-seq. Known Arabidopsis TF DNA-binding motifs were retrieved from CIS-DB version 1.02 (Weirauch et al., 2014), and those described in Franco-Zorrilla et al. (2014). Motif occurrences were determined using FIMO (Grant et al., 2011) and enrichment was assessed using the hypergeometric distribution against the background of all genes. Number of genes indicates genes containing each motif within 500 bp promoter region compared to the total number of genes in each comparison (DEG list). P-value cut off is 0.001. (XLSX)

**S8 Table.** A) Overlap between genes associated with TPR1 binding sites and genes differentially regulated in Arabidopsis plants expressing the RxL21 effector. "RxL21 induced" or "RxL21 repressed" indicates genes which show significantly higher or lower expression respectively in HA:21-expressing plants compared to HA:21ΔEAR-expressing plants. Number of genes for each comparison is shown with overlap to TPR1 targets in brackets. P values indicate significance of overlap between each group of genes with TPR1 targets. Genes selected for verification by ChIP-PCR are indicated in bold (Column D). Gene descriptions taken from Araport 11. B) The control set of 150 genes which show no differential expression between HA:21 and HA:21ΔEAR.

(XLSX)

**S9** Table. Primer sequences used in this study. (XLSX)

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