

# ATR2<sup>Cala2</sup> from Arabidopsis-infecting downy mildew requires 4 TIR-NLR immune receptors for full recognition

Dae Sung Kim<sup>1,2</sup> D, Yufei Li<sup>2</sup> D, Hee-Kyung Ahn<sup>2</sup> D, Alison Woods-Tör<sup>3</sup>, Volkan Cevik<sup>4</sup> D, Oliver J. Furzer<sup>2</sup> D, Wenbo Ma<sup>2</sup> , Mahmut Tör<sup>3</sup> and Jonathan D. G. Jones<sup>2</sup>

<sup>1</sup>State Key Laboratory of Biocatalysis and Enzyme Engineering, Hubei University, Wuhan, 430062, China; <sup>2</sup>The Sainsbury Laboratory, Norwich Research Park, Norwich, NR4 7UH, UK;

Authors for correspondence: Jonathan D. G. Jones Email: jonathan.jones@tsl.ac.uk

Mahmut Tör Email: m.tor@worc.ac.uk

Received: 11 March 2024 Accepted: 17 April 2024

New Phytologist (2024) 243: 330-344 doi: 10.1111/nph.19790

Key words: Arabidopsis, ATR2, Hyaloperonospora arabidopsidis, plant immunity, RPP2, RxLR effector, TNL.

# **Summary**

- Arabidopsis Col-0 RPP2A and RPP2B confer recognition of Arabidopsis downy mildew (Hyaloperonospora arabidopsidis [Hpa]) isolate Cala2, but the identity of the recognized ATR2<sup>Cala2</sup> effector was unknown.
- To reveal ATR2<sup>Cala2</sup>, an F<sub>2</sub> population was generated from a cross between Hpa-Cala2 and Hpa-Noks1. We identified ATR2<sup>Cala2</sup> as a non-canonical RxLR-type effector that carries a signal peptide, a dEER motif, and WY domains but no RxLR motif. Recognition of ATR2<sup>Cala2</sup> and its effector function were verified by biolistic bombardment, ectopic expression and Hpa
- ATR2<sup>Cala2</sup> is recognized in accession Col-0 but not in Ler-0 in which RPP2A and RPP2B are absent. In ATR2<sup>Emoy2</sup> and ATR2<sup>Noks1</sup> alleles, a frameshift results in an early stop codon. RPP2A and RPP2B are essential for the recognition of ATR2<sup>Cala2</sup>. Stable and transient expression of ATR2<sup>Cala2</sup> under 35S promoter in Arabidopsis and Nicotiana benthamiana enhances disease
- Two additional Col-0 TIR-NLR (TNL) genes (RPP2C and RPP2D) adjacent to RPP2A and RPP2B are quantitatively required for full resistance to Hpa-Cala2. We compared RPP2 haplotypes in multiple Arabidopsis accessions and showed that all four genes are present in all ATR2<sup>Cala2</sup>-recognizing accessions.

### Introduction

Plants, like animals, are constantly exposed to potentially damaging pathogens, and like invertebrates but unlike mammals, rely solely on innate immunity (Jones & Takemoto, 2004). The plant immune response is highly effective but must be activated early to thwart pathogens, and activation requires detection of pathogen molecules by cell surface and intracellular immune receptors. Cell-surface receptors usually detect relatively conserved pathogen-associated molecular patterns and activate pattern-triggered immunity (PTI) (Monaghan & Zipfel, 2012; Boutrot & Zipfel, 2017). During plant-microbe co-evolution, pathogens evolved the ability to deliver effector proteins to host cells that suppress PTI, enabling pathogen growth (Feng & Zhou, 2012). In turn, plants evolved intracellular immune receptors, often encoded by resistance (R) genes, that either directly or indirectly detect the presence of pathogen effector proteins (Nürnberger et al., 2004; Chisholm et al., 2006; Jones & Dangl, 2006; Jones et al., 2016) and activate effector-triggered immunity (Dodds & Rathjen, 2010; Dangl et al., 2013). According to the gene-for-gene model, resistance is determined by matching pairs of plant R-genes and pathogen avirulence (Avr)

genes. Recognized effectors are often referred to as Avr proteins. Intracellular recognition usually requires nucleotide-binding, leucine-rich repeat (NB-LRR or NLR) immune receptors. NLR activation results in an elevated immune response, characterized by generation of reactive oxygen species, cell wall fortification, activation of defence-associated genes and a localized cell death known as the hypersensitive response (HR) (Spoel & Dong, 2012). Many cases of matching R and Avr genes have been described (Bernoux et al., 2011; Césari et al., 2014; Ma et al., 2020; Redkar et al., 2023). However, in some examples, disease resistance against a pathogen isolate or recognition of an Avr protein requires the coordinate function of pairs of NLR genes (Eitas & Dangl, 2010). Recent detailed studies on the Arabidopsis TIR-NLR pair RRS1 and RPS4, and the rice coiled-coil (CC)-NLR pairs RGA4/RGA5 and Pik-1/Pik-2 reveal how such protein pairs function together. The paired partners interact physically to form a receptor complex in which each protein plays distinct roles in effector recognition or signalling activation, exemplifying a conserved mode of action of NLR pairs in diverse plants (Césari et al., 2014; Sarris et al., 2015; Ma et al., 2018). Such gene pairs are often divergently transcribed. Interestingly, 10 of 11 pairs of Toll interleukin-1 receptor (TIR)-NLR genes

<sup>&</sup>lt;sup>3</sup>Department of Biological Sciences, School of Science and the Environment, University of Worcester, Worcester, WR2 6AJ, UK; <sup>4</sup>Department of Life Sciences, The Milner Centre for Evolution, University of Bath, Bath, BA2 7AY, UK

show a head-to-head configuration in *Arabidopsis* (Meyers *et al.*, 2003; Saucet *et al.*, 2021). Divergent transcription may assure balanced levels of the protein pair to meet a strict stoichiometric requirement to act together, possibly in a complex (Narusaka *et al.*, 2009). However, the *Arabidopsis RPP2* locus that confers resistance to downy mildew (Sinapidou *et al.*, 2004) comprises two genes, *RPP2A* and *RPP2B*, which are not divergently transcribed.

In Arabidopsis, rice and many other plants, canonical and non-canonical R-Avr interactions have been found in response to a variety of microbial pathogens. Large-scale sequencing, genomics and rapid gene isolation techniques have accelerated the isolation of race-specific R-genes and their corresponding Avr (recognized effector) genes in Triticeae, including wheat, barley, rye and their wild relatives. Non-canonical NLR protein-coding genes with unique domain architectures have been found controlling race-specific resistance (Sánchez-Martín & Keller, 2021). Genetic studies of the flax-flax rust interaction identified four R-gene loci in flax that encode TIR-NLR and corresponding recognized effector genes, which encode small-secreted proteins with no locus similarity and no close homologues (Ellis et al., 2007). The world's third-largest food crop, potato, severely suffers from late blight caused by oomycete, Phytophthora infestans. This oomycete secretes host-translocated RxLR effectors, some of which are avirulence factors that are recognized by NLR proteins from wild Solanum species (Vleeshouwers et al., 2011; Lin et al., 2023).

Downy mildews are obligate biotrophic oomycete pathogens that can cause significant economic impacts on crop and ornamental plants (Thines & Kamoun, 2010; Tör et al., 2023). Downy mildew disease control is dependent on integrated strategies, incorporating cultural practices, deployment of resistant cultivars, crop rotation, systemic pesticides and biopesticides. However, downy mildews deploy large repertoires of effectors, including the so-called RxLR proteins that promote virulence or are recognized as avirulence factors (Baxter et al., 2010; Tör et al., 2023). Hyaloperonospora brassicae causes severe disease in Chinese cabbage (Brassica rapa L. ssp. pekinensis), which is native to China and is one of the most important vegetables in Asia. In epidemic seasons with warm temperatures and high humidity, 80-90% of Chinese cabbage plants are infected by H. brassicae, leading to a 30-50% reduction in production (Li et al., 2011). Downy mildew caused by Bremia lactucae is the most important disease in lettuce (Lactuca sativa L.) reducing yield and decreasing the quality of the marketable portion (Parra et al., 2021). Downy mildew caused by Plasmopara viticola can lead to severe damage to grapevines (Li et al., 2015). Cucumber (Cucumis sativus L.) downy mildew, caused by Pseudoperonospora cubensis, is a major destructive and widespread disease of cucumber plants (Zhang et al., 2019). There has been an increasing interest in the molecular mechanisms of downy mildew resistance (Liu et al., 2021). The use of cultivars carrying dominant resistant (Dm) genes in lettuce is the most effective way to control downy mildew caused by B. lactucae (Parra et al., 2021). The model plant Arabidopsis is susceptible to the downy mildew Hyaloperonospora arabidopsidis (Hpa) (Slusarenko & Schlaich, 2003). Various Resistance to *Peronospora parasitica* (*RPP*; the former name of *Hpa*) genes in different accessions confer resistance to specific *Hpa* isolates (Asai *et al.*, 2018).

Obligate biotrophic pathogen races or isolates differ in their capacity to evade or suppress host recognition (Oliver & Ipcho, 2004). Many oomycete pathogens deploy effector proteins with a signal peptide and typically the signature amino acid motifs RxLR and DEER (Rehmany et al., 2005). A subset of such effectors also carries a variable number of repeats of a WY domain (Win et al., 2012). RxLR effectors have been intensively investigated since their discovery (Anderson et al., 2015; Wood et al., 2020). RxLR genes are abundant in the genomes of *Phytophthora* and downy mildew species. Annotations based on canonical RxLR motifs predict that *Phytophthora* genomes typically contain several hundred RxLR genes (Anderson et al., 2015).

The Arabidopsis/Hpa pathosystem reveals extensive genetic diversity in host Resistance (RPP) and cognate pathogen ATR (Arabidopsis thaliana recognized) genes, which encode Hpa avirulence factors (Coates & Beynon, 2010; Asai et al., 2018). Using an Hpa reference genome (Baxter et al., 2010), 475 Hpa gene models were identified that encode effector candidates in Hpa-Emoy2, using the following criteria: (1) proteins with a signal peptide and canonical RxLR motif, such as ATR1, ATR13 and ATR39 (HaRxLs) (Allen et al., 2004; Rehmany et al., 2005; Goritschnig et al., 2012), (2) RxLR-like proteins with at least one non-canonical feature, such as ATR5 (HaRxLLs) (Bailey et al., 2011), (3) putative Crinkler-like proteins with RxLR motif (HaRxLCRNs) (Win et al., 2007) and (4) homologous proteins based on amino acid sequence similarity over the N-terminal region including a signal peptide and RxLR motif (e.g. HaRxL1b, HaRxLL2b and HaRxLCRN3b) (Asai et al., 2014).

Several RPP genes, including RPP1, RPP2A and RPP2B, RPP4, RPP5, RPP8, RPP13 and RPP39 encode NLR immune receptors (Holub, 2008). Genetic analyses of avirulence in Hpa has confirmed a gene-for-gene relationship for ATR genes (Holub et al., 1994) with their corresponding RPP genes. Recognized Hpa effectors ATR1, AvrRPP4, ATR5, ATR13 and ATR39 have been identified for RPP1, RPP4, RPP5, RPP13 and RPP39 (Allen et al., 2004; Rehmany et al., 2005; Bailey et al., 2011; Goritschnig et al., 2012; Asai et al., 2018). For example, the RPP1 locus, which contains a complex resistance gene cluster, was originally identified in Arabidopsis accession Wassilewskija (Ws-2) (Botella et al., 1998). Several members of the RPP1 gene family confer resistance against isolates of Hpa (Botella et al., 1998; Rehmany et al., 2005; Sohn et al., 2007) including RPP1-WsA, RPP1-WsB, RPP1-WsC and RPP1-NdA, while RPP1-like genes from other accessions have been implicated in hybrid incompatibility (Bomblies et al., 2007). Proteins encoded by two RPP1 alleles have been shown to recognize the cognate effector ATR1 from Hpa (Rehmany et al., 2005; Krasileva et al., 2010; Ma et al., 2020). The R proteins RPP1-WsB and RPP1-NdA share a common TNL domain architecture and are 87% identical at the amino acid level. Although polymorphisms are present throughout their coding sequences, most of the differences occur in the LRR region and include both single amino acid polymorphisms and short insertions and deletions. ATR1 belongs

to a simple locus in *Hpa* with allelic variants present in different pathogen races (Rehmany et al., 2005; Krasileva et al., 2010). ATR1 carries an N-terminal eukaryotic signal peptide and an RxLR motif (Rehmany et al., 2005; Birch et al., 2006) and associates with its cognate RPP1 immune receptor via its LRR domain (Krasileva et al., 2010). The tetrameric complex containing four RPP1 and four ATR1 molecules is mediated by direct binding of ATR1 to a C-terminal jelly-roll/Ig-like domain (C-JID) and the LRRs of RPP1 (Ma et al., 2020).

RPP2A and RPP2B are both required for resistance to Hpa isolate Cala2 (Sinapidou et al., 2004), but their cognate effector ATR2 was not identified previously. Adjacent to RPP2A (At4g19500) and RPP2B (At4g19510) (Sinapidou et al., 2004) lie two other TNL encoding genes (At4g19520 and At4g19530, hereafter RPP2C and RPP2D). They comprise, in a head-to-head conformation, a similar gene pair to RRS1 and RPS4, including a C-terminal-extended post-LRR domain. In this research, we aimed to clone and characterize ATR2 and investigate its virulence function and its contribution to effector recognition by the four genes at the RPP2 locus.

Using an F<sub>2</sub> population generated from a cross between Hpa-Cala2 and Hpa-Noks1 (Bailey et al., 2011), we positionally cloned ATR2. We show here that functional ATR2 is absent from the reference Emoy2 genome and its annotated proteome, that ATR2 confers elevated disease susceptibility when expressed in planta and that all four RPP2 paralogs contribute to its full recognition.

### **Materials and Methods**

# Plant materials and growth

Arabidopsis accessions, Col-0, Ler-0, Oy-0, Ws-2, Ws-2 eds1 and CW84, which is an *Hpa*-susceptible recombinant inbred line generated from a cross between Col-0 and Ws-2 (Botella et al., 1998) were grown at 22°C under short-day condition (10 h : 14 h, light : dark) and Nicotiana benthamiana plants were grown at 25°C under a 16 h : 8 h, light : dark period in environmentally controlled growth cabinets.

# Positional cloning of ATR2<sup>Cala2</sup>

The crossing of Hpa-Cala2 and Hpa-Noks1 and production of F<sub>2</sub> mapping population from a single-spored CaNo F1 were described previously (Bailey et al., 2011). Initially, segregating 52 random CaNo F<sub>2</sub> isolates were bulked up on Ws-eds1 seedlings and tested on Col-5 to determine the genetic nature of ATR2. As the genomic sequences of parental isolates were not available then, a similar approach to clone ATR5 (Bailey et al., 2011) was taken where DNA was isolated from individual CaNo F2 isolates and a bulk segregant analysis was employed to clone ATR2. Two different bulks were constructed from the CaNo F2 individuals (18 F2s with ATR2/± and 17 with atr2/atr2 genotypes), and AFLP was carried out with EcoRI and MseI primer pairs as described (Bailey et al., 2011). Fifteen polymorphic AFLP fragments were

identified and converted to CAPS markers to map ATR2 onto publicly available BAC contigs. As the Hpa-Emoy2 reference genome became available, we used these markers to identify the Hpa-Emoy2 SuperContig9. As the number of recombinants were very low, additional CaNo F2 isolates were generated, and Illumina paired-end sequencing data of CaNo F2 bulks were obtained. As the genomic data for Hpa-Cala2 and Hpa-Noks1 became available (Woods-Tör et al., 2018), the bulk sequences were mapped onto Hpa-Cala2 genome as described (Woods-Tör et al., 2018) and SNP markers were identified within the interval. Further markers were generated from the identified SNP sites, and using a total of 130 CaNo F<sub>2</sub> isolates, we mapped ATR2 to a 186.5 kb interval on Hpa-Cala2 SuperContig9. Further markers were generated and new F<sub>2</sub> isolates were obtained, and the locus was mapped to a 112 kb interval. We compared genomic sequences of Hpa-Emoy2, Hpa-Noks1and Hpa-Cala2 for the interval to identify possible candidates for ATR2. All PCR amplifications for mapping were performed as described (Woods-Tör et al., 2018).

# Pathogen assays

Hpa isolates, Hpa-Emoy2, Hpa-Noks1 and Hpa-Cala2, were propagated and maintained by weekly sub-culture on 14-d-old Arabidopsis seedlings. Preparation of inoculum for experiments and the assessment of sporulation were as described previously in Bailey et al. (2011).

Pseudomonas syringae pv tomato (Pst) DC3000 was grown in King's B broth (10 g peptone, 15 g glycerol, 1.5 g K<sub>2</sub>HPO<sub>4</sub> and 5 mM MgSO<sub>4</sub> per litre) containing 50 μg ml<sup>-1</sup> rifampicin. Leaves of 5-wk-old Arabidopsis plants were infiltrated with 10<sup>5</sup> CFU ml<sup>-1</sup> of Pst DC3000 using a needleless syringe. Bacterial growth was measured at 0- and 3-d post inoculation (dpi).

Phytophthora infestans isolate 88069 was grown on Rye Agar at 19°C for 2 wk. Plates were flooded with 5 ml of cold H<sub>2</sub>O and scraped with a glass rod to release zoospores. The resulting solution was collected in a falcon tube and zoospore numbers were counted using a hemacytometer and adjusted to  $2 \times 10^4$  zoospores ml<sup>-1</sup> and 10  $\mu$ l droplets were inoculated onto the abaxial side of leaves of intact N. benthamiana plants. Inoculated leaves were then stored on moist tissues in sealed boxes.

### Plasmid construction

All the constructs used in this study were generated using Uracil-Specific Excision Reagent (USER) enzyme cloning method (Geu-Flores et al., 2007). Briefly, target DNA to be cloned into destination USER vectors, pICSLUS0003 or pICSLUS0004 (archived in The Sainsbury Laboratory) was amplified using  $PfuTurbo^{\mathbb{R}}$   $C_x$ polymerase (Agilent Technologies, Santa Clara, CA, USA) with uracil-containing primer pair then assembled with desired tag ('Hellfire' including 6-His and 3-FLAG epitopes), linearized vector and USER enzyme (NEB, Ipswich, MA, USA). For transient gene expression in N. benthamiana or Nicotiana tabacum, ATR2 candidates without signal peptide were cloned and assembled.

# Bombardment and luciferase assays

Co-bombardment assays were performed as described previously with some modifications (Bailey *et al.*, 2011). Briefly, *Arabidopsis* plants were grown with short-day condition until 6 wk old. Detached leaves were placed on a 1% MS agar in a Petri dish. One micrometre of tungsten particles was coated with the plasmids carrying genes *ATR2* and luciferase under 35S promoter. Bombardments were performed using a Bio-Rad PDS-1000 (He) apparatus with 1100 p.s.i. rupture discs, as per the manufacturer's instructions. For each replicate, a leaf from both test and control plant genotypes were co-bombarded together in a single shot. Bombarded leaves were put into 10-ml plastic vials filled with water 1 cm from the bottom and were incubated at 25°C for 20 h.

For the luciferase assay, a Dual Reporter Luciferase Assay system (Promega, Madison, WI, USA) was used. Four transiently bombarded leaf events were pooled together and crushed in Luciferase Cell Culture Lysis buffer (Promega). The extract was centrifuged at 11 269 *g*-force for 10 min at 4°C. Twenty microlitres of the lysate was then dispersed in 96-well plates in triplicates and analysed on Varioskan Flash Instrument (Thermo Fisher Scientific, Waltham, MA, USA) by injecting 100 µl of luciferase assay reagent II, which includes substrate and reaction buffer. A 10-s read time was used to measure luciferase activity for each well.

# **Expression analysis**

Total RNA was isolated from three biological replicates using the RNeasy Plant Mini Kit (Qiagen, Hilden, Germany) with the Dnase treatment (Qiagen). cDNA was synthesised using Super-Script IV Reverse Transcriptase (Thermo Fisher Scientific). For *ATR2* gene expression analysis during *Hpa* infection, reverse transcription polymerase chain reaction (RT-PCR) was performed.

# Transient expression in Nicotiana species

Agrobacterium tumefaciens GV3101 strain harbouring ATR2 candidate fused to 35S promoter was streaked on selective media and incubated at 28°C for 24 h. A single colony from the streaked inoculum was transferred to liquid LB media with appropriate antibiotic and incubated at 28°C for 48 h in a shaking incubator at 5 g-force. The cultures were centrifuged at 2012 g-force for 5 min and resuspended in infiltration buffer (10 mM MgCl<sub>2</sub>, 10 mM MES, pH 5.7), and acetosyringone was added to a final concentration of 200 μM at OD<sub>600</sub> of 1.0. The abaxial surface of 4-wk-old N. tabacum or N. benthamiana was infiltrated with a 1-ml needleless syringe (Kim et al., 2015).

# Arabidopsis transformation

Arabidopsis accessions Col-0 and Ler-0 expressing ATR2 candidate gene, and CW84 expressing Col-RPP2 cluster harbouring JAtY clone (Zhou et al., 2011), were transformed using A. tume-faciens strain GV3101 by flower dipping method (Clough & Bent, 1998).

# RPP2 cluster haplotype analyses

Full-length amino acid sequences of individual RPP2A, RPP2B, RPP2C and RPP2D from 64 different *Arabidopsis* accessions were extracted from pan-NLRome data (Van de Weyer *et al.*, 2019). Each group of RPP2 was aligned to each other using the GENEIOUS PRIME software to investigate haplotype patterns of RPP2 clusters. PFAM (Punta *et al.*, 2012; http://pfam-legacy.xfam.org) was used for domain analysis in RPP2 cluster.

# Protein structure modelling

Protein tertiary structure model of full-length ATR2<sup>Cala2</sup> was generated by Alphafold 2 (Jumper *et al.*, 2021; Varadi *et al.*, 2021). The region spanning the Y-WY sequences was extracted and superimposed with the structure of full-length PsPSR2 using PyMOL Molecular Graphics System, v.1.2r3pre, LLC (Xiong *et al.*, 2014; He *et al.*, 2019; Hou *et al.*, 2019). Secondary structures and surface accessibility of ATR2<sup>Cala2</sup> were predicted by NetSurfP-3.0 (Høie *et al.*, 2022). Alignment with published LWY effectors revealed the conserved W and Y residues in ATR2<sup>Cala2</sup> and the corresponding Y and WY modules (He *et al.*, 2019).

### Accessions

Genomic sequences of parental isolates can be found under accession nos. GCA\_001414265.1 for *Hpa*-Cala2, GCA\_001414 525.1 for *Hpa*-Noks1 and GCA\_000173235.2 for *Hpa*-Emoy2 in NCBI. The raw sequence reads from the genomics sequencing of bulks are available from the Sequenced Read Archive (SRA) under accession nos. SRX13788375 (avirulent) and SRX1 3788374 (virulent). *ATR2*<sup>Cala2</sup> and *ATR2*<sup>Emoy2</sup> sequences were deposited in NCBI (GenBank accession nos. ON994189 and ON994190, respectively). Resistance (*R*) gene sequence capture (RenSeq) raw sequencing data of FN2 (*rpp2a-1*) mutant is available from the SRA (accession no. PRJNA955397).

Supporting Information Methods S1 contains additional methods, including protein gel blot, R gene sequence capture (RenSeq), and bioinformatics.

### Results

# Mapping ATR2

Positional cloning was used to identify the ATR2 locus in Hpa-Cala2. A segregating CaNo  $F_2$  population (Bailey  $et\ al.,\ 2011$ ) was used to define the ATR2 locus. Initially, 52 randomly chosen  $F_2$  isolates were tested on Col-5. A single semi-dominant avirulence determinant designated  $ATR2^{Cala2}$  segregated in the  $F_2$  population (avirulence: virulence ratio was 40:12, with chi-square = 0.1025 and P=0.74; Table S1). Bulked segregant analysis was used to identify AFLP markers that are linked to ATR2 in the CaNo  $F_2$  population. AFLP markers were cloned and converted to CAPS markers, which were then used for mapping ATR2. The reference genome of Hpa-Emoy2 was still being

generated during this early mapping work, and genomic sequence data for Hpa-Cala2 and Hpa-Noks1 were not available. Once Hpa-Emoy2 genomic sequence became available, we transferred the AFLP-derived CAPS markers to *Hpa*-Emoy2 SuperContig9, which helped us to identify the physical location of the locus (Figs \$1, \$2).

We then generated 100 bp paired-end Illumina HiSeq2500 sequencing data from the two newly bulked (virulent and avirulent) pools, comprising 110 million reads for the virulent bulk and 104 million reads for the avirulent bulk. We also utilized Hpa-Cala2 and Hpa-Noks1 genomic sequences (Woods-Tör et al., 2018) to identify SNPs between Hpa-Cala2 and Hpa-Noks1 genomes. Using new markers generated from these SNPs, we established an interval of 186.5 kb on Hpa-Emoy2 Super-Contig9: 656515-843042 (Fig. S1). We generated further markers and F2 isolates and narrowed the ATR2 locus to a 112 kb interval on Hpa-Emoy2 SuperContig9: 708503-820527 (Tables 1, S2). Genes in the ATR2 interval encode effector-like proteins (Results S1; Figs S2–S4).

# A2C3<sup>Cala2</sup> encodes an RxLR effector-like candidate for ATR2<sup>Cala2</sup>

A2C3 was identified in Hpa-Cala2 after re-sequencing of the 5 kb upstream of A2C1 and A2C2, which includes a highly polymorphic region of Cala2 compared with Emoy2 (Fig. 1a). We found a transposable element in this region of the *Hpa*-Emoy2

genome and a 2.3 kb deletion in Hpa-Cala2. We also found a cytosine insertion on the Hpa-802071 coding region in Cala2, which created a frameshift in the Hpa-802071 coding region (Fig. 1b). To determine whether A2C3<sup>Cala2</sup> co-segregates with recognition by RPP2 in the F2 population, A2C3 alleles were amplified and sequenced from 12 CaNo F2 segregants, Hpa-Emoy2, Hpa-Noks1 and Hpa-Cala2 (Fig. S5). All avirulent F2s were homozygous or heterozygous for A2C3<sup>Cala2</sup>, while all virulent  $F_2$ s were homozygous for  $A2C3^{Emoy2/Noks1}$  (Figs 1c, S5c). Of Hpa-Emoy2, Hpa-Noks1 and Hpa-Cala2, only Hpa-Cala2 carries this non-canonical RxLR effector candidate (A2C3<sup>Cala2</sup>), which has a signal peptide, a dEER, and Y and WY motifs (Figs 2a, S6a).  $A2C3^{Emoy2}$  and  $A2C3^{Noks1}$  alleles were identical to each other with early stop codons caused by a frameshift. These data suggested A2C3<sup>Cala2</sup> might be ATR2<sup>Cala2</sup>. We analysed synonymous and non-synonymous SNPs in A2C3 alleles among seven different *Hpa* isolates for which genomic data are available. Ten non-synonymous SNPs are found only in Hpa-Cala2, indicating specificity of A2C3<sup>Cala2</sup> (Table S3). Alignment of A2C3<sup>Cala2</sup> with *Phytophthora* LWY effectors revealed conserved W and Y residues and the corresponding Y and WY modules of A2C3<sup>Cala2</sup> (He et al., 2019). The Y-WY modules of A2C3<sup>Cala2</sup> in an A2C3<sup>Cala2</sup> structural model predicted by Alphafold 2 were extracted and superimposed with PsPSR2, which is a typical RxLR effector with one WY motif and six LWY motifs. This region was well-matched on PsPSR2 Y5-LWY6 (region from Y5 of LWY5 to LWY6) with Root Mean Square Deviation

Table 1 Intervals of ATR2 from CaNo F2 isolates. Red: Cala2 homozygote; Yellow: Noks1 homozygote; Orange: Cala2-Noks1 heterozygote at A2C3.

Isolate F2	Marker <sup>a</sup> 656515	708503	759570	801023	820527	826062	843042	Col-0 Phenotype <sup>b</sup>
Cala2	Cala2°	Cala2	Cala2	Cala2	Cala2	Cala2	Cala2	Av
Noks1	Noks1 <sup>d</sup>	Noks1	Noks1	Noks1	Noks1	Noks1	Noks1	V
2	Noks1/Cala2°						Noks1/Cala2	Av
9	Noks1/Cala2						Noks1/Cala2	Av
10	Noks1/Cala2						Noks1/Cala2	Av
15	Noks1/Cala2						Noks1/Cala2	Av
21	Cala2						Cala2	Av
30	Cala2	Noks1/Cala2	Noks1/Cala2	Noks1/Cala2	Noks1/Cala2	Noks1/Cala2	Noks1	Av
34	Noks1/Cala2	Cala2	Cala2	Cala2			Cala2	Av
45	Noks1	Cala2	Cala2	Cala2			Cala2	Av
104	Noks1		· · · · · · · · · · · · · · · · · · ·				Noks1	V
113	Noks1	Noks1	Noks1	Noks1	Noks1	Noks1	Noks1/Cala2	V
118	Noks1						Noks1	V

<sup>&</sup>lt;sup>a</sup>Each marker number indicates nucleotide no. on SuperContig9 from Emoy2 genomic sequence as a reference.

eCala2-Noks1 heterozygote.



<sup>&</sup>lt;sup>b</sup>Virulent (V) or avirulent (Av) on Col-0.

Cala2 homozygote.

<sup>&</sup>lt;sup>d</sup>Noks1 homozygote.

Marker

746974

CACC**C**GGT

CACC-GGT

CACC**C**GGT

CACC**C**GGT

CACC**C**GGT

CACC**C**GGT

CACC-GGT

CACC-GGT

CACC-GGT

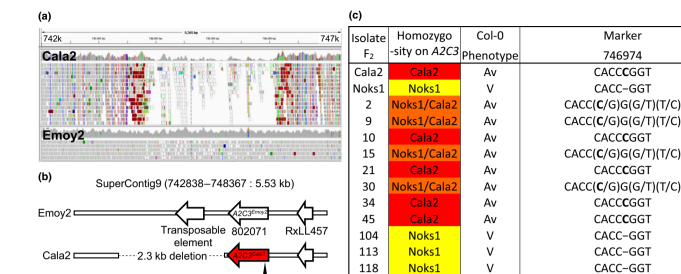


Fig. 1 Genetic determination of  $A2C3^{Cala2}$  co-segregation from avirulent  $F_2$  isolates. (a) Comparison of polymorphic region of Cala2 with Emoy2 from 742k to 747k of SuperContig9. Screenshot was captured from the Icv software. (b) Transposable element next to '802071' on Emoy2 and 2.3 kb deletion on Cala2 of assigned region. A2C3<sup>Cala2</sup> allele is highlighted with red colour. A cytosine (C) insertion at 746974 indicated with a black triangle. (c) Analyses of homo- or heterozygosity, avirulent (Av) or virulent (V) on Col-0 and a segregated cytosine (C) insertion in Av isolates at the frameshift region (746974) by sequencing from Supporting Information Fig. \$5. Red: Cala2 homozygote; Yellow: Noks1 homozygote; Orange: Cala2-Noks1 heterozygote at A2C3. Inserted cytosine was bold highlighted.

(RMSD) = 2.305 (Fig. 2b). This structural comparison also revealed that there is an L-like module that harbours several leucines (L) contributing to the additional hydrophobic core formation between Y and WY modules of A2C3<sup>Cala2</sup> even though this was not predicted by amino acid sequence comparison (Figs 2b, S6b).

To reveal whether A2C3<sup>Cala2</sup> has weighted sequence similarity to other RxLR proteins in the post-SP and pre-WY region, a Hidden Markov Model was used using 475 published Hpa effectors including RxLR, RxLL and CRN effectors. No aligned sequences of A2C3<sup>Cala2</sup> were found likely due to the diversification of N-terminal sequences of Hpa effectors. Therefore, we constructed a phylogenetic tree using post-SP N-terminal sequences of 476 Hpa effectors including A2C3<sup>Cala2</sup> and found only a few effectors cluster in a clade with A2C3<sup>Cala2</sup>. Even ATR5, another non-RxLR effector, is not related with A2C3<sup>Cala2</sup>, and previously defined ATR effectors are distributed throughout the tree indicating there is no specific signature in N-terminal sequences of Hpa effectors (Fig. **S7**).

We determined the expression of A2C3<sup>Cala2</sup> alleles during Hpa infection. Arabidopsis Oy-0 accession was infected with Hpa-Emoy2 and Ler-0 was infected with Hpa-Cala2, and Col-eds1 was used as hyper-susceptible control.  $A2C3^{Cala2}$  is expressed at 3 dpi (Fig. S8a). Previously, Asai et al. (2014) performed expression profiling of Hpa genes from Hpa Emoy2. RNA-seq data of Hpa-802071 ( $A2C3^{Emoy2/Noks1}$ ) were retrieved from the data. Again, A2C3<sup>Emoy2/Noks1</sup> is induced after infection and shows the highest expression at 3 dpi (Fig. S8b). We proceeded to further evaluate the A2C3<sup>Cala2</sup> allele as a strong candidate for ATR2<sup>Cala2</sup>.

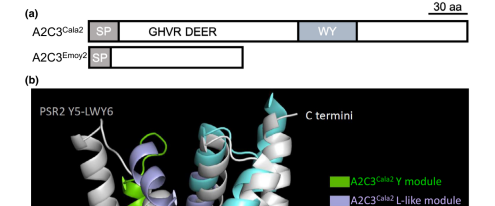
# A2C3<sup>Cala2</sup> triggers defence in Col-0

When luciferase assays were performed to evaluate the recognition of A2C3<sup>Cala2</sup> in Arabidopsis, a reduction of at least fivefold in luciferase activity was detected in Col-0 compared with empty vector (EV) control. Equal luciferase activity was detected in an Hpa-susceptible recombinant inbred line Arabidopsis CW84 when leaf tissue was bombarded with 35S:A2C3<sup>Cala2</sup> or EV. These results suggest that Col-0 but not CW84 can recognize A2C3<sup>Cala2</sup>. As before, CCG28 was recognized by WRR4A, which served as a positive control (Fig. 2c) (Redkar et al., 2023). Thus, our genetic investigations and bombardment experiments are consistent with A2C3<sup>Cala2</sup> being the avirulence determinant ATR2<sup>Cala2</sup>, and we hence refer to A2C3<sup>Cala2</sup> as ATR2<sup>Cala2</sup>. As an additional test of ATR2<sup>Cala2</sup> detection by RPP2 in *Arabidopsis*, ATR2<sup>Cala2</sup> under 35S promoter was transformed into Col-0. Only three T<sub>1</sub> lines were selected from antibiotic screening, and strikingly, all three transformants showed strong dwarf phenotype, consistent with recognition of ATR2<sup>Cala2</sup> in Arabidopsis Col-0 background (Fig. 2d).

# ATR2<sup>Cala2</sup> enhances susceptibility in the absence of host recognition

Plant pathogen effector proteins that are translocated into host cells can attenuate host defence. Many pathogen effectors interfere with cellular processes that are essential for innate immunity.

To evaluate its virulence function, ATR2<sup>Cala2</sup> was transiently expressed in N. benthamiana leaves that were then inoculated with P. infestans race 88069. The P. infestans lesion area was N termini



A2C3<sup>Cala2</sup> WY module

RMSD = 2.305

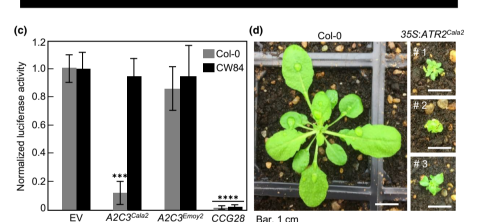


Fig. 2 A2C3<sup>Cala2</sup> recognition capacity in Col-0. (a) Schematic diagrams of A2C3 of Hpa Cala2 and Emoy2. (b) Alphafold 2 prediction of Y and WY modules of A2C3<sup>Cala2</sup>, and super-imposition with PsPSR2. PsPSR2 contains seven (L)WY units with Y5-LWY6 showing the highest similarity with A2C3<sup>Cala2</sup>. This structural comparison also revealed an 'L'-like fold between the 'Y' and 'WY' sequence in A2C3<sup>Cala2</sup>. (c) Biolistic bombardment of A2C3 with luciferase into Col-0 and CW84 in which RPP2 is absent. CCG28 recognized by WRR4A in Col-0 and CW84 served as a positive control. Data are mean  $\pm$  SD from three independent experiments. Asterisks (\*\*\*, P < 0.001; \*\*\*\*, P < 0.0001) indicate statistical significance compared with luciferase alone in Col-0 or CW84 by two-way ANOVA with Tukev's multiple comparison test. EV, empty vector. (d) Transgenic Arabidopsis Col-0 expressing ATR2<sup>Cala2</sup> under 35S promoter. Photographs were taken from the 4-wk-old plants under short-day condition. Bars, 1 cm.

significantly larger in  $ATR2^{Cala2}$ -expressing leaf sectors than in GFP vector control (Fig. 3a). At 7 dpi, lesion area in the  $ATR2^{Cala2}$ -expressing region was more than four times larger than that observed in GFP control region (Fig. 3b). Stable  $ATR2^{Cala2}$ -expressing Arabidopsis lines (35S: $ATR2^{Cala2}$ ) were generated in Ler-0, which lacks RPP2A and RPP2B. In contrast to  $ATR2^{Cala2}$  expressing Col-0, all transgenic lines selected grew similar to Ler-0 wild-type (WT; Fig. 3c). Strikingly, all the transgenic lines were more susceptible to virulent Pst DC3000 or Hpa-Cala2 compared with Ler-0 WT control (Fig. 3d,e). Ler-eds1 was used as hyper-susceptible control. Collectively, these data show that in both Arabidopsis and N. benthamiana,  $ATR2^{Cala2}$  expression can compromise plant innate immunity in the absence of recognition by a cognate R-gene.

In addition to RPP2A and RPP2B, two additional linked TNLs, RPP2C and RPP2D, are required for full RPP2 function

We tested the requirement for RPP2A (At4g19500) and RPP2B (At4g19510) in ATR2<sup>Cala2</sup> recognition. There are two other

adjacent TIR-NB-LRR genes (At4g19520 and At4g19530, hereafter RPP2C and RPP2D) (Fig. S9a) that comprise a gene pair similar to RRS1 and RPS4, with C-terminal-extended post-LRR domains and a head-to-head orientation (Fig. S9a,b). RPP2A contains two TIR-NB-ARC domains connected by the Arabidopsis LSH1 and Oryza G1 (ALOG) domain followed by LRRs towards its C terminus. Post-LRR (PL) domains of RPP2B and RPP2D are homologous to the RPP1 C-terminal jelly-roll/Ig-like (C-JID) domain. RPP2C harbours an additional TIR domain following an extended post-LRR domain (Fig. S9b; Table S4). We obtained the fast neutron 2 (FN2) rpp2a mutant (Sinapidou et al., 2004), and several T-DNA insertional mutants from GABI or SALK for rpp2b, rpp2c and rpp2d (Fig. S9c). We combined sequence capture with Illumina sequencing (RenSeq) with DNA from the FN2 (rpp2a-1) mutant and confirmed a 25 bp deletion in RPP2A. The RPP2B, RPP2C and RPP2D mutations were also verified (Fig. S10). After inoculating mutants with Hpa-Cala2, conidiospores were counted at 7 dpi. Ler-0 and Ws-eds1 were used as susceptible controls. While fewer than  $1 \times 10^2$  spores per plant were detected in the resistant Col-0, c.  $4 \times 10^3$  spores per plant were detected on rpp2a-1 and rpp2b-1 mutants with

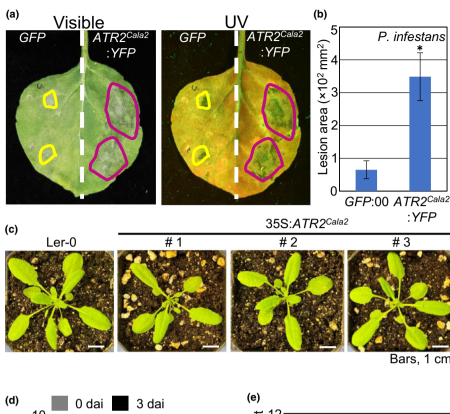
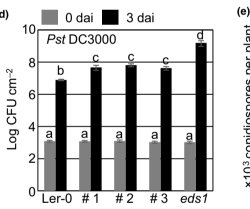
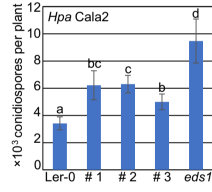


Fig. 3 Enhanced disease susceptibility resulting from exogenous ATR2<sup>Cala2</sup> expression. (a) Phenotypes on Nicotiana benthamiana transiently expressing the GFP control (GFP) or ATR2<sup>Cala2</sup>:YFP under the 35S promotor followed by Phytophthora infestans 88069 inoculation  $(2 \times 10^4 \text{ zoospores ml}^{-1})$  2 d after transient expression. Photographs were taken 7 days after inoculation (dai) with P. infestans. (b) Disease lesion area of P. infestans on N. benthamiana leaves. Forty lesion squares of each were measured. (c) Generation of constitutively ATR2<sup>Cala2</sup> expressing transgenic Arabidopsis in Ler-0 background. Bars, 1 cm. (d) Bacterial growth in Ler-0, ATR2<sup>Cala2</sup>-OX Ler-0 (# 1, # 2 and #3) and Ws-eds1 (eds1) as a hypersusceptible control infected with Pst DC3000  $(10^5 \text{ CFU ml}^{-1})$ . (e) Quantification of conidiospores on wild-type and transgenic plants at 7 dai infected with Hpa Cala2  $(5 \times 10^4 \text{ conidiospores ml}^{-1})$ . Data are means  $\pm$  SD from three independent experiments. Asterisks indicate significant differences as determined by Student's *t*-test (P < 0.05). According to Fisher's Least Significant Difference (P < 0.05), statistical significance was shown by different letters above each bar.

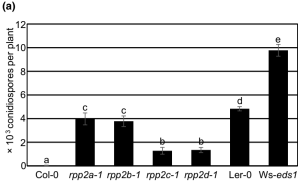


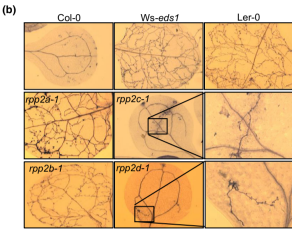


similar values to those obtained from Ler-0 (near  $4.8 \times 10^3$  per plant), indicating Cala2 resistance in Col-0 is compromised by rpp2a or rpp2b mutations (Fig. 4a). Interestingly, rpp2c-1 and rpp2d-1 mutants also showed compromised resistance to Hpa-Cala2. Around  $1.2 \times 10^3$  to  $1.3 \times 10^3$  spores per plant were counted from rpp2c and rpp2d mutants, suggesting RPP2C and RPP2D also contribute to full resistance against Hpa-Cala2 in Col-0 (Fig. 4a). Trailing necrosis was observed on rpp2c and rpp2d mutants, while no necrosis was observed on Col-0 at 6 dpi (Fig. S11a). To visualize cell death and hyphal growth, we performed trypan blue staining at 5 dpi using infected cotyledons. Local cell death was observed on Col-0, and hyphal growth and haustoria formation over the whole leaf was observed on rpp2a, rpp2b and Ler-0 cotyledons, as well as Ws-eds1. Partial but restricted hyphal growth was detected on rpp2c and rpp2d mutants (Fig. 4b).

Transgenic complementation assays with CW84 were carried out using JAtY 49E17 clone (Zhou et al., 2011), which harbours the whole RPP2 cluster. While sporangiophore formation was observed on CW84, complemented transgenic plants restored complete resistance to *Hpa*-Cala2 (Fig. S11b). When *Hpa*-Emoy2 was inoculated onto *rpp2a*, *rpp2b*, *rpp2c* and *rpp2d* mutants, resistance was not compromised, due to *RPP4*-dependent resistance in Col-0 (van der Biezen et al., 2002; Fig. S12), which is why the susceptible phenotypes were only seen after inoculation with *Hpa*-Cala2.

To assess  $ATR2^{Cala2}$  recognition capacity by RPP2 paralogs, luciferase eclipse assays were conducted using individual Col-0 rpp2a-1, rpp2b-1, rpp2c-1 and rpp2d-1 mutants. The luciferase activity was normalized to compare with that of EV control on Col-0 (Fig. 4c). The normalized luciferase activity in each individual Col-0 rpp2a mutant and Col-0 with EV was comparable





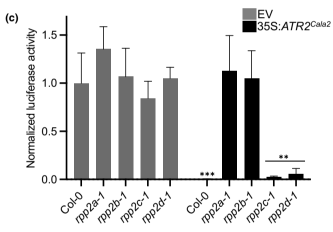


Fig. 4 Compromised Hpa Cala2 resistance in rpp2 mutants. (a) Quantification of conidiospores on Col-0, individual rpp2 mutants from Col-0, Ler-0 and Ws-eds1 at 7 days after inoculation (dai) infected with Hpa Cala2 (5  $\times$  10<sup>4</sup> conidiospores ml<sup>-1</sup>). Data are means  $\pm$  SD from three independent experiments. According to Fisher's Least Significant Difference, (P < 0.05), statistical significance was shown by different letters above each bar. (b) Trypan blue staining of Hyaloperonospora~arabidopsidis~hyphal~growth~on~cotyledons~at~5~dai.~Hyphal~growth~region~on~rpp2c~and~rpp2d~mutants~was~enlarged~to~clearly~show~the~Hpa~hyphal~development. (c) Luciferase~measurement~upon~biolistic~bombardment~into~Col-0~and~rpp2~mutants.~Statistical significance~compared~with~luciferase~alone~in~Col-0~is~indicated~by~asterisks~(\*\*\*, <math>P < 0.01; \*\*\*\*, P < 0.001) according to two-way~ANOVA~with~Tukey's~multiple~comparison~test.

with no significant differences indicating particle bombardment distributed well through the leaves of Col-0 and each mutant. When  $ATR2^{Cala2}$  was bombarded together with 35S:luciferase on

Col-0, normalized luciferase activity was strongly reduced, ranging from 0.004 to 0.007, while those on rpp2a-1 or rpp2b-1 still maintained a range of 0.86–1.55, indicating ATR2<sup>Cala2</sup> recognition is almost completely abolished in rpp2a-1 and rpp2b-1 mutants. ATR2<sup>Cala2</sup> was still recognized in rpp2c-1 and rpp2d-1 mutants, with normalized activity ranging from 0.01 to 0.1 (Fig. 4c). Even though no statistically significant differences were detected between  $ATR2^{Cala2}$ -bombarded Col-0, rpp2c-1 and rpp2d-1, the average values of the luciferase activities on rpp2c-1 (mean, 0.024) and rpp2d-1 (mean, 0.054) are almost 5–10 times higher than on Col-0 (mean, 0.005) when co-bombarded with  $ATR2^{Cala2}$ , consistent with RPP2C and RPP2D weakly contributing to ATR2<sup>Cala2</sup> recognition. These data indicate RPP2C and RPP2D are also required for full ATR2<sup>Cala2</sup>-triggered immunity.

To monitor whether the co-expression of RPP2A, RPP2B, RPP2C and RPP2D with or without ATR2Cala2 can trigger HR, we transiently expressed RPP2A, RPP2B, RPP2C or RPP2D, either alone or co-expressing RPP2A/RPP2B, RPP2C/RPP2D, RPP2A/RPP2B/RPP2C/RPP2D or RPP2A/RPP2B/RPP2C/RPP 2D in combination with ATR2<sup>Cala2</sup> in N. benthamiana. Rpiamr3 and Avramr3, an R-gene from Solanum americanum and the recognized P. infestans RxLR Avr gene, respectively, were used as a HR-inducing positive control (Lin et al., 2022). No auto-activity of any RPP combination was observed, either by individual RPP2 paralogs or by co-expression of RPP2 paralogs. Similarly, no HR was observed when all four RPP2 were expressed in combination with ATR2<sup>Cala2</sup> (Fig. S13). We performed co-immunoprecipitation (co-IP) to monitor whether ATR2<sup>Cala2</sup> can interact with each RPP2 protein using N. benthamiana transient expression assays. However, under our experimental conditions, no association was detected between ATR2<sup>Cala2</sup> and any RPP2 protein (Fig. S14). RPP2C protein was not detected by Western blot; we infer that more detailed work might be required to correctly define its splicing and thus where to attach an epitope tag to its C terminus. However, RPP2C expression was abundantly detected by reverse transcription polymerase chain reaction indicating that the TAIR annotation that we used to tag RPP2C (RPP2C.1) is probably wrong and did not enable us to express the protein in transient assays. Alternatively, RPP2C might be differentially spliced between Arabidopsis and N. benthamiana (Fig. S15).

# RPP2 haplotype diversity

As the *RPP2* cluster containing *RPP2A*, *RPP2B*, *RPP2C* and *RPP2D* is required for full ATR2<sup>Cala2</sup>-triggered resistance, we assessed *RPP2* haplotype diversity in multiple *Arabidopsis* accessions. An investigation of the *Arabidopsis* pan-NLRome (Van de Weyer *et al.*, 2019) enabled in-depth analysis for the *RPP2* cluster. We compared the *RPP2* cluster in 64 *A. thaliana* accessions (Fig. S16). Col-0, Oy-0 and Can-0 have the complete form of the *RPP2* cluster, while other accessions lack some *RPP* genes or harbour incomplete (partial) *RPP* genes (Fig. 5a). Interestingly, almost all accessions contain a complete form of *RPP2B*, and seven ecotypes among 21 harbour *RPP2A* while many of other ecotypes have incomplete alleles of *RPP2A* (Fig. 5a). Almost half

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**Fig. 5** Differential  $ATR2^{Cala2}$  recognition capacity dependent on RPP2A haplotype. (a) Heatmap diagram for *RPP2* cluster haplotype analyses from 21 Arabidopsis ecotypes. (b) Normalized luciferase activity by biolistic bombardment of  $ATR2^{Cala2}$  with luciferase into Col-0, Ler-0 (*RPP2A*, *2B*-lacking) and Ws-2 (partial *RPP2A*). Data are means  $\pm$  SD from three independent experiments. Asterisk indicates a significant difference as determined by two-way ANOVA with Tukey's test (\*\*\*\*, P < 0.0001). (c) RPP2A haplotype analyses from 21 Arabidopsis ecotypes.

of accessions lack, or contain partial forms of, *RPP2C* or *RPP2D* (Fig. 5a). Among 64 accessions, while only 17 accessions contain complete *RPP2A* encoding TIR-NB-TIR-NB-LRR (TN-TNL) homologues, the *RPP2B*-encoding TIR-NB-LRR (TNL) is well-conserved in almost all accessions excluding Ler-0, Rsch-4 and Vig-1 (Figs S16, S17a). *RPP2C* is lacking or incomplete in > 40 accessions and the amino acid length of RPP2D is quite diverse (Fig. S16). While RPP2A haplotypes show structural diversity on their first TIR-NB-ARC domains, RPP2B, RPP2C and RPP2D are more conserved in different *Arabidopsis* accessions (Fig. S17).

To monitor ATR2<sup>Cala2</sup> recognition capacity, luciferase eclipse assay bombardment was conducted with Col-0 (complete RPP2 cluster), Ler-0 (*RPP2C* and *RPP2D* lacking) and Ws-2 (partial *RPP2A* and *RPP2C* lacking). As expected, ATR2<sup>Cala2</sup> is recognized in Col-0, while Ler-0 and Ws-2 lack ATR2<sup>Cala2</sup> recognition capacity, indicating a critical role of *RPP2A* and *RPP2B* for ATR2<sup>Cala2</sup> recognition (Fig. 5c). Compared with other *RPP2* genes, *RPP2A* was present in diverse forms. As shown in Fig. 5 (c), diverse accessions including Ws-2 lost the first N-terminal TIR-NB-ARC domain. The TIR-NB-ARC defect in Ws-2 abolishes ATR2<sup>Cala2</sup> recognition.

# **Discussion**

Downy mildews such as *B. lactucae* on lettuce (Parra et al., 2021), *P. viticola* on grapevines (Li et al., 2015), *P. cubensis* on cucumber (Zhang et al., 2019) and *H. brassicae* on brassicas (Liu et al., 2021) are destructive obligate oomycete phytopathogens on fruit and vegetable crops (Thines & Kamoun, 2010; Tör et al., 2023). Genetic variation for downy mildew resistance has also been studied in *Brassica* species such as *B. napus*, broccoli, non-heading Chinese cabbage and Chinese cabbage (Chen et al., 2008; Xiao et al., 2016). The 27 known *Dm* genes in lettuce are located in gene clusters that encode NLRs (Parra et al., 2021). A better understanding of resistance mechanisms to downy mildew is highly desirable. *Arabidopsis* NLR-encoding *RPP* genes confer recognition of specific downy mildew races and different RPP proteins specifically recognize their cognate downy mildew RxLR effectors (Asai et al., 2018).

Most oomycete pathogens deploy secreted effector proteins, with the signature amino acid motif RxLR, which enter plant cells where they promote virulence (Win et al., 2012; Asai et al., 2014; Wood et al., 2020). The function and evolution of RxLR effectors have been investigated since their discovery (Anderson et al., 2015). Comparative genomics indicates that RxLR genes play a major role in virulence for downy mildews and Phytophthora species. Although progress has been made, there is still much to learn about the mechanisms of downy mildew virulence and host resistance. Most P. infestans and Hpa effectors carry an RxLR motif.

We positionally identified  $ATR2^{Cala2}$  that encodes a non-canonical RxLR-like protein recognized by RPP2A and RPP2B.  $ATR2^{Cala2}$  encodes an RxLR-like protein with an N-terminal signal peptide, and dEER and C-terminal Y and WY modules. The ATR2 alleles in other Hpa strains are identical to the  $ATR2^{Emoy2}$  allele and lack GHVR, dEER and WY motifs due to a frame shift

resulting from a single nucleotide deletion. In the absence of recognition by RPP2A and RPP2B, ATR2<sup>Cala2</sup> expression enhances pathogen susceptibility in planta. Furthermore, the head-to-head RPP2C and RPP2D genes, which are adjacent to RPP2A and RPP2B, also contribute to full resistance to Hpa-Cala2. ATR5 was the first example of a non-canonical RxLR effector lacking the canonical RxLR motif but with an Nterminal signal peptide and a canonical EER motif (Bailey et al., 2011). At the expected RxLR position, ATR5 carries Gly-Arg-Val-Arg (GRVR) instead of RxLR. ATR2<sup>Cala2</sup> at this position carries Gly-His-Val-Arg (GHVR) followed by a dEER motif. ATR5 contains two WY motifs and one LWY motif at its C terminus (Fig. S18a). The Y-WY domain of ATR2<sup>Cala2</sup> resembles the LWY of ATR5 based on Alphafold2 structural prediction (Fig. S18b). In the Hpa genome, > 150 genes encode for potentially secreted proteins such as ATR2<sup>Cala2</sup> that carry motifs such as signal peptide and EER but lack the RxLR motif (Asai et al., 2014). This also been seen in other oomycetes such as Pseudoperonospora and Bremia (Purayannur et al., 2020; Wood et al., 2020; Nur et al., 2023). We conclude that although the RxLR motif is often found in oomycete effectors, in Hpa as in other oomycetes, some divergence is permitted for effector translocation. Therefore, additional Hpa effectors may exist that have not yet been predicted.

RPP2 was the first genetically defined R-gene locus shown to carry two NLR-encoding genes, both of which are required for function (Sinapidou et al., 2004). The corresponding recognized effector from Hpa enables investigations into how the RPP2-encoded immune receptor complex functions. Recognized effectors are also valuable tools for investigating plant-microbe interactions, since their host targets correspond to important plant defence components. Most R-gene pair-encoding NLR proteins, such as Arabidopsis RRS1-RPS4 that recognize bacterial effectors AvrRps4 and PopP2, and rice RGA4-RGA5 recognizing rice blast effectors AVR1-CO39 and AVR-Pia, are encoded by divergently transcribed genes (Césari et al., 2013; Ma et al., 2018), in contrast to RPP2A and RPP2B (Fig. S9a). Sensor NLRs are dependent on executor (or helper) NLRs for downstream immune signalling (Feehan et al., 2020). RRS1 functions as a sensor that reveals effectors that target WRKY domain transcription factors, while RPS4 is an executor (Ma et al., 2018). Uniquely, RPP2A contains two TIR-NB-ARC domains followed by LRR, and an ALOG domain that is specific and conserved to land plants and has DNA-binding activity (Yoshida et al., 2009; Naramoto et al., 2020; Beretta et al., 2023) between the two TIR-NB-ARC domains (Fig. S9). Comparative analyses of RPP2A in diverse Arabidopsis accessions show that the main variation in the RPP2A haplotype is the presence or absence of one N-terminal TIR-NB-ARC (Figs 5c, S16). RPP2B is a typical TIR-NLR resembling the executor NLR, RPS4. Compared with RPP2A, RPP2B is relatively well-conserved in different Arabidopsis accessions (Figs S16, S17a). Conceivably, RPP2A functions as a sensor for ATR2<sup>Cala2</sup> and RPP2B functions as a signal executor. TIR domains of plant NLRs are known to have nicotinamide adenine dinucleotide hydrolase (NADase) activity, which requires a catalytic glutamate (E), that activates defence (Wan

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et al., 2019). The C-JID domains of Arabidopsis RPP1 and N. benthamiana ROQ1 (recognition of XopQ1) are required for pathogen effector recognition. ATR1 binds to the C-JID and the LRRs of RPP1 leading to assembly of tetramers with NADase activity (Ma et al., 2020). The LRR and C-JID of ROQ1 directly interact with Xanthomonas effector (XopQ), allowing the NB-ARC domain to transition to an ATP-bound state. Complex assembly results in TIR proximity that opens the NADase active site (Martin et al., 2020). The first TIR on RPP2A has the catalytic E residue, whereas the second TIR on RPP2A lacks the conserved E residue (Table S4). The TIR of RPP2B has the conserved E residue and a C-JID (Table S4), consistent with a role as executor. Still, the domains of RPP2A and/or RPP2B that interact with ATR2<sup>Cala2</sup> remain to be elucidated.

We also revealed the requirement for two additional TIR-NB-LRR genes, RPP2C and RPP2D, adjacent to RPP2A and RPP2B and showed all four NLR proteins are required for full resistance against Hpa-Cala2. A paired head-to-head R-gene structure is often found in plant-paired NLRs (Narusaka et al., 2009; Césari et al., 2014; Saucet et al., 2021). RPP2C and RPP2D form a head-to-head orientation similar to RRS1-RPS4 (Narusaka et al., 2009; Ma et al., 2018; Guo et al., 2020). The C-terminal post-LRR domain of RPS4 is homologous with C-JID suggesting that it recognizes conformational changes in RRS1 upon effector recognition (Saucet et al., 2021). The RPP2C post-LRR domain is homologous to that of RRS1 but RPP2C contains a TIR domain on its C-terminal end instead of WRKY. RPP2D contains a C-JID on its C terminus homologous to that of RPS4 and RPP1 (Fig. S9; Table S4). Many Arabidopsis accessions lack or have incomplete RPP2C but RPP2D is relatively conserved among different accessions (Fig. S16). Even though RPP2C and RPP2D are quantitatively required for full resistance against Hpa-Cala2, how this pair contributes to ATR2<sup>Cala2</sup> recognition remains unclear. Conceivably, their contribution could either be additive, or by potentiating RPP2A/B-dependent recognition. We speculate that since both RPP2A and RPP2C carry integrated TIR domains, which do not contain a catalytic E residue, ATR2<sup>Cala2</sup> might function by interacting with and somehow suppressing functions of host TIR domain-containing proteins. RPP4, a TIR-NLR, can mediate HR upon detection of Hpa RxLR effector AvrRPP4 (Asai et al., 2018). ATR2<sup>Cala2</sup> transient expression could not suppress RPP4-mediated HR (Fig. \$19). Interestingly, the TIR on RPP2C C-terminal end lacks the catalytic E residue for NADase activity, while the first TIR on RPP2C has the E residue, and RPP2D also has the E residue on its TIR (Table S4). We hypothesize that the second TIR on RPP2A might function to detect ATR2<sup>Cala2</sup> leading to conformational change via RPP2B interaction with ATR2<sup>Cala2</sup>. This could result in RPP2A/B resistosome activation enabling signal transduction through activated NADase function of the first TIR on RPP2A and RPP2B TIR. If the TIR in RPP2A lacking catalytic E functions as an effector decoy, the C-terminal TIR on RPP2C might also act as an integrated decoy to detect ATR2 Cala2. However, thus far we were unable to detect direct or indirect interaction between ATR2<sup>Cala2</sup> and each RPP2 protein (Fig. S14). Even

though *RPP2C* expression was abundantly detected by reverse transcription polymerase chain reaction (Fig. S15), no signal was detected for C-terminal epitope tagged RPP2C by Western blot (Fig. S14). Searching for the actual splicing pattern of 3' of *RPP2C* and determining its encoding amino acid sequence is required to investigate the mechanisms of an NLR complex involving four TNL proteins. Further research is needed to define the effector recognition mechanisms for these atypical NLR protein pairs.

# **Acknowledgements**

Financial support from the Gatsby Charitable Foundation (http://www.gatsby.org.uk/), and from BBSRC grants BB/K009176/1 and BB/M003809/1 to JDGJ, is gratefully acknowledged. This work is also supported in part by the grant 09 963/A from the Leverhulme Trust to MT. We thank Matthew Smoker and Jodie Taylor for their help with *Arabidopsis* transformation. The authors would like to thank Dr Kenichi Tsuda for providing luciferase assay kit for DSK.

# **Competing interests**

None declared.

# **Author contributions**

DSK, MT and JDGJ conceptualized and designed the research. DSK, AW-T, VC and OJF conducted all experiments. DSK and MT performed the data analysis. VC and MT gave critical intellectual input and provided materials for this work. YL and WM carried out structural prediction and analyses of *Hpa* effectors. H-KA carried out protein-complex analyses. DSK, MT and JDGJ wrote the manuscript with input from all co-authors.

# **ORCID**

Hee-Kyung Ahn https://orcid.org/0000-0002-8884-0156
Volkan Cevik https://orcid.org/0000-0002-3545-3179
Oliver J. Furzer https://orcid.org/0000-0002-3536-9970
Jonathan D. G. Jones https://orcid.org/0000-0002-4953-261X

Dae Sung Kim https://orcid.org/0000-0002-4579-2094
Yufei Li https://orcid.org/0000-0003-0245-3807
Wenbo Ma https://orcid.org/0000-0001-5569-639X
Mahmut Tör https://orcid.org/0000-0002-4416-5048

# Data availability

All the sequence data used in this study can be found in NCBI (see the Materials and Methods section; https://www.ncbi.nlm.nih.gov/sra/SRX13788375; https://www.ncbi.nlm.nih.gov/sra/SRX13788374; https://www.ncbi.nlm.nih.gov/nuccore/ON994189.1/; https://www.ncbi.nlm.nih.gov/nuccore/ON994190.1/; https://www.ncbi.nlm.nih.gov/bioproject/PRJNA955397/).



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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 Map-based cloning approach for ATR2<sup>Cala2</sup> allele.
- Fig. S2 Diagram of a contig in which ATR2 candidates co-segregate.
- Fig. S3 Loci, structure and recognition of A2C1 and A2C2.
- **Fig. S4** *A2C1* and *A2C2* do not trigger cell death with *RPP2* in *Nicotiana* species.
- **Fig. S5** Determination of  $A2C3^{Cala2}$  allele co-segregation on avirulent *Hyaloperonospora arabidopsidis* Cala2-Noks1 F2 population.
- Fig. S6 Protein sequence analyses of A2C3<sup>Cala2</sup>.
- **Fig. S7** Phylogenetic analysis of post-SP N-terminal sequences of 475 *Hyaloperonospora arabidopsidis* Emoy effectors and A2C3.
- **Fig. S8** A2C3 expression after *Hyaloperonospora arabidopsidis* Emoy2 and Cala2 infection.
- Fig. S9 Scheme of RPP2 cluster.
- **Fig. S10** RenSeq-MiSeq from *rpp2a* mutant. Red triangle indicates 25 bp deletion in *RPP2A*.
- Fig. S11 Compromised ATR2-mediated resistance in *rpp2* mutants.
- **Fig. S12** *Hyaloperonospora arabidopsidis* Emoy2 growth on Col-0, individual *rpp2* mutants and Oy-0.

- **Fig. S13** No auto-activity causing hypersensitive response was observed from an individual or co-expression of *RPP2* paralogs in *Nicotiana benthamiana*.
- Fig. S14 ATR2 is not associated with each RPP2 protein.
- **Fig. S15** Reverse transcription polymerase chain reaction RT-PCR analyses of *RPP2C* expression.
- Fig. S16 RPP2 cluster haplotype analyses from pan-NLRome (Van de Weyer *et al.*, 2019).
- Fig. S17 Haplotype analyses of RPP2B, RPP2C and RPP2D.
- **Fig. S18** Comparison of WY domains of ATR2<sup>Cala2</sup> and ATR5.
- **Fig. S19** No suppressive activity of ATR2 on RPP4-triggered hypersensitive response.
- Methods S1 Protein gel blot, RenSeq and bioinformatics.
- **Results S1** Genes in the *ATR2* interval encode effector-like proteins.
- **Table S1** Interaction phenotypes recorded after inoculation of Col-5 with CaNo F<sub>2</sub> isolates.
- **Table S2** Mapping table of ATR2.
- **Table \$3** Number of SNPs on A2C3 alleles on seven different *Hyaloperonospora arabidopsidis* isolates.
- **Table S4** Domains on RPP2 proteins and catalytic E residue conservation on each Toll interleukin-1 receptor.

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