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# Allelic variants of the NLR protein Rpi-chc1 differentially recognize members of the *Phytophthora infestans* PexRD12/31 effector superfamily through the leucine-rich repeat domain

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#### SUMMARY

Phytophthora infestans is a pathogenic oomycete that causes the infamous potato late blight disease. Resistance (R) genes from diverse Solanum species encode intracellular receptors that trigger effective defense responses upon the recognition of cognate RXLR avirulence (Avr) effector proteins. To deploy these R genes in a durable fashion in agriculture, we need to understand the mechanism of effector recognition and the way the pathogen evades recognition. In this study, we cloned 16 allelic variants of the Rpi-chc1 gene from Solanum chacoense and other Solanum species, and identified the cognate P. infestans RXLR effectors. These tools were used to study effector recognition and co-evolution. Functional and non-functional alleles of Rpi-chc1 encode coiled-coil nucleotide-binding leucine-rich repeat (CNL) proteins, being the first described representatives of the CNL16 family. These alleles have distinct patterns of RXLR effector recognition. While Rpi-chc1.1 recognized multiple PexRD12 (Avrchc1.1) proteins, Rpi-chc1.2 recognized multiple PexRD31 (Avrchc1.2) proteins, both belonging to the PexRD12/31 effector superfamily. Domain swaps between Rpi-chc1.1 and Rpi-chc1.2 revealed that overlapping subdomains in the leucine-rich repeat (LRR) domain are responsible for the difference in effector recognition. This study showed that Rpi-chc1.1 and Rpi-chc1.2 evolved to recognize distinct members of the same PexRD12/31 effector family via the LRR domain. The biased distribution of polymorphisms suggests that exchange of LRRs during host-pathogen co-evolution can lead to novel recognition specificities. These insights will guide future strategies to breed durable resistant varieties.

Keywords: NLR cluster, leucine-rich repeat, *Phytophthora infestans*, late blight resistance gene, RXLR effector, *Solanum* species, potato.

#### INTRODUCTION

Potato (*Solanum tuberosum*) is the third largest food crop in terms of human consumption in the world, after rice (*Oryza sativa*) and wheat (*Triticum aestivum*), with more than 370 million tonnes produced in 2019 (FAO, 2020; Devaux *et al.*, 2020). Potato late blight, caused by the oomycete *Phytophthora infestans*, is one of the most infamous potato diseases. During the mid-1840s, this pathogen caused the Great Irish Famine, from which around one million people died (Callaway, 2013). Nowadays, losses from late blight are estimated to reach 16% of the world production. The main disease management is based on biocide applications. Including yield losses and crop protection measures, late blight causes a global economic loss of  $\in$ 5.2 billion per year (Haverkort *et al.*, 2016).

Phytophthora infestans is an oomycete with sexual and asexual life cycles, which exhibits a hemibiotrophic lifestyle on potato. Together with its large and fast evolving genome (estimated to be 240 Mb), its population diversity leads to the regular emergence of new aggressive and

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virulent strains (Haas et al., 2009). The infection starts when a spore lands on the plant surface, germinates, and forms a penetration structure called appressorium. Alternatively, spores can also enter through natural openings such as stomata. After passing the epidermis, hyphae spread intercellularly projecting haustorium structures into the mesophyll cells. These haustoria are specialized infection structures that secrete both apoplastic and cytoplasmic effectors to create an intimate association with the host cell and facilitate nutrient uptake (Fry, 2008). Effectors are pathogen molecules that interact with different host targets to suppress the host defense response and enable colonization. The publication of the P. infestans T30-4 genome revealed the presence of 563 effector genes encoding the conserved Arg-any amino acid-Leu-Arg (RXLR) peptide motif (Haas et al., 2009). These effectors rapidly evolve by gaining and losing repeat-rich domains through recombination with different paralogs, transposon movement, and point mutations (Goss et al., 2013). During co-evolution, potato has evolved receptors to recognize some of these effectors and trigger an immune response.

Wild Solanum species are the main source of resistance (R) genes to P. infestans (Rpi). To date, over 20 Rpi genes have been characterized in different Solanum species, e.g., R1, R2, R3a, R3b, R8, and R9a from Solanum demissum, Rpi-blb1 and Rpi-blb12 from Solanum bulbocastanum, Rpivnt1 from Solanum venturii, and Rpi-amr1 from Solanum americanum (Ballvora et al., 2002; Huang et al., 2005; Jo et al., 2015; Li et al., 2011; Lokossou et al., 2009; Pel et al., 2009; van der Vossen et al., 2003, 2005; Vossen et al., 2016; Witek et al., 2021). All these receptors belong to the nucleotide-binding (NB)-leucine-rich repeat (NLR) type of receptors and contain a coiled-coil (CC) domain in their Ntermini, referred to as CC-NB-LRR (CNL). The recognition of a specific effector or avirulence factor (Avr) leads to the activation of effector-triggered immunity (ETI) and the restriction of the pathogen growth (Jones et al., 2016). ETI is mostly monogenic and therefore well suited and commonly deployed for resistance breeding and crop protection strategies. To keep up with the fast evolution of effectors, NLR genes are also very diverse and rapidly evolving. Gene duplications, recombinations, unequal crossing-over, and transpositions have been proposed to provide the basis for the evolution of the NLR recognition spectrum (Leister, 2004; Mcdowell and Simon, 2006). This fast evolution can lead to the independent development of new receptors in different geographical locations that recognize the same effector. For instance, the effector Avr2 from P. infestans is recognized by the unrelated R2 and Rpi-mcq1 CNLs (Aguilera-Galvez et al., 2018). R2 is located on chromosome IV in the Mexican species S. demissum, while *Rpi-mcq1* is located on chromosome IX from a Peruvian accession of Solanum mochiquense (Foster et al., 2009; Smilde et al., 2005). More recently, Rpi-amr1 alleles

have been described to cause differential recognition of Avramr1 homologs from several *P. infestans* isolates (Witek *et al.*, 2021). When the doubled-monoploid DM1-3 519 R44 potato genome was published, 755 NLR genes were identified (Jupe *et al.*, 2013). Many of them were found in clusters together with closely related paralogs. All of these clusters were formed in ancestral species and had sequence homology to syntenic genomic regions from other *Solanum* species harboring late blight resistance genes. Thus, alleles from functional *Rpi* genes that do not provide resistance (*rpi*) can be found in all studied *Solanum* genomes.

Here, we studied Solanum chacoense, a diploid wild potato relative from South America considered a source of resistance to P. infestans (Karki et al., 2021; Vossen et al., 2009). We identified two functionally distinct receptors, Rpi-chc1.1 and Rpi-chc1.2, which are allelic variants that recognize distinct P. infestans effectors from the same PexRD12/31 effector superfamily. Remarkably, only Rpichc1.1 is able to provide resistance against several P. infestans isolates. The expression and recognition of PexRD12 effectors was associated with Rpi-chc1.1-mediated resistance and, therefore, they were designated as Avrchc1.1 effectors. PexRD31 effectors were still expressed in several *P. infestans* isolates, but were rapidly downregulated during the interaction with potato. This potentially explains the inability of *Rpi-chc1.2* to provide late blight resistance. We postulate that *Rpi-chc1.2* is a ubiquitous ancient *R* gene that was recently overcome and PexRD31 may have functioned as Avrchc1.2. An allele-mining strategy revealed Rpi-chc1 orthologs in different wild Solanum accessions and potato cultivars that could be classified by their sequence and recognition spectrum of Avrchc1.1 or Avrchc1.2 or non-functionality. Finally, using domain swaps, we found that the LRR domain harbors the recognition specificity of both Avrchc1.1 and Avrchc1.2. The specificities resided in overlapping LRR subdomains and could not be combined into one active protein using domain exchanges.

## RESULTS

#### Cloning and characterization of Rpi-chc1.1

The *S. chacoense* accession CHC543 from Bolivia is a previously described wild potato relative harboring resistance to *P. infestans* (Vleeshouwers *et al.*, 2011a). To identify the genetic locus of resistance, the resistant seedling CHC543-5 was crossed with the susceptible seedling CHC544-5 to generate the F1 population 7650, consisting initially of 212 individuals. This population was challenged with *P. infestans* isolate 90128 in a detached leaf assay (DLA). A clear 1:1 segregation was observed, indicating the presence of a single dominant resistance gene which will henceforth be referred to as *Rpi-chc1*. Cleaved amplified polymorphic

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sequence markers from chromosome 10 were tested as this chromosome was known to harbor *Rpi-ber* from the related species *Solanum berthaultii* (Vossen *et al.*, 2013). The marker TG63 in chromosome 10 was indeed linked to the *Rpi-chc1* resistance. Successive fine-mapping in a recombinant population representing 2357 individuals was performed using markers derived from RH89-39-16 BAC clones from chromosome 10 (PGSC) (Sharma *et al.*, 2013; The Potato Genome Sequencing Consortium, 2011). A narrow genetic window between markers RH106G03-T and RH97D21\_C21-4 was identified to contain *Rpi-chc1* (Figure 1a). To generate a physical map of the mapping interval, two bacterial artificial chromosome (BAC) clones, B1 and B2, were selected from a BAC library that was derived from CHC543-5 genomic DNA. After sequencing the BAC clones, two and six NLR genes were identified in clones B1



Figure 1. Map-based cloning of Rpi-chc1.1.

(a) Genetic map of *P. infestans* (isolate 90128) resistance from CHC543-5. The number between the markers represents the number of recombinants found in a population derived from 2357 seedlings. Markers starting with RH were derived from BAC end sequences generated by PGSC. Marker B2-S represents the BAC end marker from clone 2. The black horizontal line represents the interval of *Rpi-chc1.1*. (b) Two BAC clones were isolated to generate the physical map. Annotation revealed the presence of NB-LRR genes, genes with or without predicted function, and transposable elements. Three complete NB-LRR (B2-1, B2-2, and B2-3) genes between flanking markers RH106G03T and B2-S were selected as candidates. (c) The three candidates were expressed through agroinfiltration in *N. benthamiana* leaves. An empty vector (EV) and *Rpi-blb1* were used as negative and positive controls, respectively. Only candidate B2-3 was able to compromise the growth of *P. infestans* isolate 90128. (d) The three candidates were stably transformed into the potato variety Desiree. After inoculation with isolate 90128, only the candidate B2-3 was able to provide resistance. Untransformed Desiree and Desiree plants stably transformed with *Rpi-blb1* were used as negative and positive controls, respectively.

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and B2, respectively. Further fine-mapping revealed that only the last six were located within the mapping interval and only three (B2-1, B2-2, and B2-3) encoded complete NLR proteins (Figure 1b). The three candidates were subcloned including their native 5' and 3' regulatory elements, and complementation analyses were performed in Nicotiana benthamiana. After 2 days, the agroinfiltrated area was challenged with P. infestans 90128. Rpi-blb1, which was shown to provide resistance to P. infestans, was used as a positive control. The leaves agroinfiltrated with candidate B2-3 and Rpi-blb1 showed severely compromised pathogen growth, while leaves with candidates B2-1 and B2-2 were completely susceptible to P. infestans 90128 (Figure 1c). This result suggested that B2-3 is the gene in CHC543-5 that provides resistance to *P. infestans*. To verify this result, the three candidates B2-1, B2-2, and B2-3 were stably transformed into the susceptible S. tuberosum cv. 'Desiree'. Indeed, only the events containing candidate B2-3 showed resistance to P. infestans (Figure 1d). Furthermore, three different single guide RNAs (sgRNAs) were designed to specifically target the LRR domain of

candidate B2-3 (Figure 2a). The resistant CHC543-5 genotype was stably transformed with CRISPR-Cas9 and these sgRNAs. The transformation events were challenged with *P. infestans* 90128 and IPO-C isolates, and 48% of the transformants had become susceptible to both isolates (Table S1; Figure 2b), which suggested that the active late blight resistance gene was specifically and successfully mutated. Four transformants were genotyped and frameshift mutations were found only in the three susceptible transformants (Figure 2c). Therefore, we concluded that B2-3 is the gene from CHC543-5 that is causal for late blight resistance. Henceforth, we refer to gene B2-3 as *Rpichc1.1* as it is the first *Rpi-chc1* allele that is identified in *S. chacoense*.

*Rpi-chc1.1* encodes a CNL protein that belongs to the CNL16 (Witek *et al.*, 2016) immune receptor family (Figure 3a). *Rpi-chc1.1* has one uninterrupted open reading frame of 3909 bp, which is predicted to be translated into 1303 amino acids (Figure S2). No introns were predicted. The CC domain contains the N-terminal MADA motif, four predicted  $\alpha$ -helices, and the typical hhGRExE, but the



Figure 2. CRISPR-Cas9-induced frameshifts in candidate B2-3 (*Rpi-chc1.1*) lead to *Phytophthora* susceptibility. (a) Three different sgRNAs were designed to target the LRR domain of candidate B2-3. (b) A construct with Cas9 and all three sgRNAs was stably transformed into the resistant genotype CHC543-5. Transformation events were inoculated with *P. infestans* 90128 and IPO-C isolates. 48% of the transformation events became susceptible to both isolates (Table S1). (c) One resistant (#1) and three susceptible events (#21, #24, #26) were genotyped together with the untransformed plant CHC543-5. The three susceptible events contained frameshift mutations, while the tested resistant event had no mutations in the B2-3 candidate.



Figure 3. Rpi-chc1 allele mining.

(a) The *Rpi-chc1.1* allele belongs to the immune receptor family. Different motifs were found in the different CNL receptor domains. The LRR domain consists of 29 imperfect repeats. (b) Sixteen *Rpi-chc1.1*-like sequences were cloned from eleven different diploid *Solanum* accessions. The phylogenetic analysis of the DNA sequences led to the identification of four clades. The branch lengths represent the percentage of phylogenetic distance. In the protein alignment mismatches are highlighted in red, gaps are indicated by dashes (–), and insertions relative to Rpi-chc1.1 (anchor) are indicated by a blue bracket.

distinctive EDVID motif was less conserved (Rairdan *et al.*, 2008). The NB domain contains the characteristic Kinase 1a VYND, Kinase 2, and Kinase 3a motifs (Campbell, 2003; Leipe *et al.*, 2004; Pal *et al.*, 2007; Saraste *et al.*, 1990; Wendler *et al.*, 2012). The ARC1 domain contains the RNBS-C motif, Motif 3, and the GLPL motif. The ARC2 domain contains Motif 2, the RNBS-D motif, and two copies of the MHDL motif (Danot *et al.*, 2009; Reubold *et al.*, 2011; Sukarta *et al.*, 2016). The LRR domain consists of 29 imperfect repeats. Both LRR3 and LRR4 contain a central VLDL motif which is conserved in the third LRR of most functional NLRs (Bendahmane *et al.*, 2002; Warren *et al.*, 1998).

#### Identification of Rpi-chc1.1 allelic variants

In order to identify different *Rpi-chc1.1* allelic variants, we pursued an allele-mining approach in closely related resistant and susceptible *S. chacoense, S. berthaultii, Solanum tarijense,* and *S. tuberosum* accessions. Homologous sequences were amplified using primers overlapping with the start and stop codons of *Rpi-chc1.1*. The PCR fragments of the expected 3.9 kb size were cloned and sequenced, resulting in the identification of 15 *Rpi-chc1.1*-like sequences. The mined *Rpi-chc1.1* variants contained between 1296 and 1303 amino acids (Figure 3b; Figure S2). From the selected diploid accessions one or two sequence variants were identified, suggesting that indeed *Rpi-chc1* alleles were mined rather than paralogs. In case only one variant was mined from an accession, this suggested that

the second allele has significant sequence polymorphisms at (one of) the primer annealing sites. Phylogenetic analysis of the sequences showed strong sequence similarities among the alleles (94.6-100% identity). Even within this high identity range, the presence of four main clades was revealed (Figure 3b). In clade 1, the Rpi-chc1.1 allele was found, together with three sequences from S. berthaultii that were nearly identical to each other and a sequence from S. tarijense. From clade 1, together with Rpi-chc1.1, we selected one sequence from S. berthaultii (94-2031) and one from S. tarijense (TAR852-5) for complementation analysis. Transformation of the corresponding genes to susceptible Desiree plants showed that they provide resistance to P. infestans isolates 90128 and IPO-C, like Rpichc1.1 (Table S2). We therefore concluded that clade 1 contains functional homologs of Rpi-chc1.1. The S. tarijense allele will be referred to as Rpi-tar1.1. The S. berthaultii allele will be referred to as Rpi-ber1.1, which matches to the previously described Rpi-ber and Rpi-ber1 genes that were derived from the same accession (PI473331) at similar genetic positions (Rauscher et al., 2006; Tan et al., 2010; Vossen et al., 2013).

The allele mining in accession CHC543-5 resulted not only in the re-identification of the active *Rpi-chc1.1*, but also in the identification of a presumed allelic variant, which we will refer to as *Rpi-chc1.2*. To test if *Rpi-chc1.1* and *Rpi-chc1.2* were indeed alleles of the same gene, we tested *Rpi-chc1.2*-specific markers in the recombinant population 6750 (CHC543-5  $\times$  CHC544-5). We found a perfect

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repulsion between Rpi-chc1.2 and Rpi-chc1.1, strongly suggesting that both genes are allelic variants (Table S3). Additionally, this analysis proved that Rpi-chc1.2 does not cause resistance against P. infestans 90128, even though Rpi-chc1.2 is expressed during infection (Table S2; Figure S1f). The Rpi-chc1.2 protein sequence clusters in clade 2 together with four identical sequences from S. berthaultii. Close to clade 2, we can observe clade 3, which consisted of an S. berthaultii, an S. tarijense, and an S. tuberosum allele from RH89-039-16, a diploid clone previously characterized as susceptible to P. infestans (Vleeshouwers et al., 2011a). The clade 3 allele from S. tari*jense* contained an in-frame stop codon, making it unlikely that this allele produces an active resistance protein. Clade 4 contained only S. berthaultii alleles. The allelic variants were numbered according to the clade in which they were found (i.e., Rpi-ber1.1 from clade 1, Rpi-ber1.2 from clade 2, etc.), followed by an extension to indicate the genotype from which the allele was derived.

## Rpi-chc1.1 recognizes the RXLR PexRD12 effector family from *P. infestans*

To understand the resistance mechanism of the *S. chacoense* CHC543-5 accession, we searched for the effector recognized by Rpi-chc1.1. A collection of 90 *P. infestans* extracellular (Pex) proteins in a PVX agroinfectious vector, of which 54 contained the RXLR-DEER motif (PexRD), was screened. Individual clones from the Pex collection were co-agroinfiltrated with *Rpi-chc1.1* in *N. benthamiana* leaves. As a positive control, we used a mix of the *R3a*/ *Avr3a R* gene effector pair, which is known to trigger a strong hypersensitive response (HR) in *N. benthamiana* leaves. Only two effectors from the Pex collection, PexRD12-1 and PexRD12-2 (PITG\_16233 and PITG\_16240, respectively), were able to trigger an *Rpi-chc1.1*-dependent HR (Figure 4a). Neither the inactive paralogs B2-1 and B2-2 nor R3a produced an HR upon co-agroinfiltration with PexRD12. These results showed that PexRD12 is specifically recognized by Rpi-chc1.1. We could further confirm this finding using transgenic Desiree potato plants that were transformed with Rpi-chc1.1. About half of this transgenic population showed late blight resistance, while the other half was susceptible, probably due to impaired transgene expression. Interestingly, the plants that showed late blight resistance also showed PexRD12 recognition, while the susceptible transgenic plants did not show any response upon PexRD12 agroinfiltration (Table S4). We sought for further evidence that PexRD12 was indeed causing avirulence on Rpi-chc1.1-expressing plants. In a field trial with natural infection, we found isolates that were virulent on plants containing Rpi-chc1.1. The infected material was collected and used for gene expression analysis via quantitative reverse transcription-PCR (RT-qPCR). PexRD12 expression was not detected in the Rpi-chc1.1 resistance-breaking isolates, while other effector genes such as Avrsto1 were normally expressed (Vleeshouwers et al., 2008). Reciprocally, we found that Rpi-sto1 breaking isolates still expressed PexRD12 normally (Figure 4b). Altogether, these results suggest that PexRD12 corresponds to Avrchc1.1.

# The PexRD12/31 superfamily is a complex *P. infestans* RXLR effector family

Using Blast analyses of the T30-4 proteome, we found nine homologs of PexRD12 in the *P. infestans* T30-4 genome. Additionally, we found that PexRD12 proteins had strong homology with nine members of the PexRD31 family and two additional, more distantly related sequences (Table S5). These 20 effectors are encoded by clusters of paralogs mainly in three supercontigs (Figure S3) and will henceforth be referred to as the PexRD12/31 superfamily (see also Petre *et al.*, 2020). All PexRD12/31 effectors are





(a) The three *Rpi-chc1.1* candidates were co-agroinfiltrated with the Pex effector collection in *N. benthamiana* leaves to screen for Avrchc1.1. Rpi-chc1.1 induces cell death when co-expressed with both PITG\_16233 and PITG\_16240, from the PexRD12 family. *R3a* and *Avr3a* were used as negative controls, and a mix of *R3a* and *Avr3a* was used as a positive control. (b) Relative *Avrchc1.1* and *Avrsto1* effector expression was measured in infected plant material from a field trial with natural *P. infestans* infection (2013, Wageningen). Untransformed Desiree plants and Desiree plants transformed with *Rpi-chc1.1* (Desiree:*Rpi-chc1.1*) or *Rpi-sto1* (Desiree:*Rpi-sto1*) were used in this study. Relative expression data were obtained by dividing the relative effector gene expression by the expression of the *P. infestans* elongation factor 2 (*ef2*) gene. Three independent samples were included in the RT-qPCR experiment. Stars represent a statistically significant difference (two-sample *t*-test, *P* < 0.008).

small proteins that include a signal peptide in the Nterminus, an effector domain in the C-terminus, and the conserved RXLR and EER motifs in the center, except for PITG\_16243 and PITG\_09577, which contain RXXR-EER and RXXLR-EER motifs, respectively (Figure 5a).

The alignment of the protein sequences and the phylogenetic analysis of the PexRD12/31 superfamily members resulted in five main clades (Figure 5a). Two highly homologous clades, PexRD12-A1 and PexRD12-A2, can be distinguished to form the PexRD12 family. Clade PexRD12-A2 also includes truncated versions which partly or completely miss the effector domain. In addition, two related clades, PexRD31-B and PexRD31-C, constitute the PexRD31 family. Additionally, PITG\_16428 and PITG\_09577 were much less related and are together referred to as PexRD12/ 31 group D.

To determine the degree to which PexRD12/31 members are expressed *in planta*, as observed for other Avr effectors of *P. infestans* (Rietman *et al.*, 2012; Vleeshouwers *et al.*, 2011b), we tested their expression during infection with qPCR on cDNA using clade A-, B-, and C-specific primers. The relative expression was calculated and normalized for

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the relative amount of P. infestans. Three different P. infestans isolates were evaluated at different time points after inoculation of different susceptible potato genotypes (Figure S1a-d). In all the tested genotypes, PexRD12 showed the highest relative expression. In two isolates maximum expression was found from 4 to 24 h after inoculation and expression remained high until after 48 h in all four isolates. The PexRD31-B effectors were expressed in two isolates but were rapidly downregulated in the first hours after inoculation with hardly any expression left. PexRD31-C expression was mostly undetectable along the inoculation time course. Similar results were observed when PexRD12/31 expression was analyzed during infection of the four different P. infestans isolates EU\_13\_A2, Ec1, EU\_6\_A1, and US23 (Figure S4), using the data from the PenSeg dataset (Lin et al., 2020).

# Rpi-chc1.2 recognizes the RXLR PexRD31 effector family from *P. infestans*

In order to describe the spectrum of effector recognition by different *Rpi-chc1* alleles, several representatives from each clade were selected and co-agroinfiltrated with



#### Figure 5. Rpi-chc1 alleles show non-overlapping recognition of the PexRD12/31 effector superfamily.

(a) Twenty members of the PexRD12/31 superfamily were found in *P. infestans* isolate T30-4. In the amino acid sequence, we can distinguish a signal peptide in the N-terminus, the conserved RXLR-EER motifs in the center, and the effector domain in the C-terminus. Some PexRD12/31 family members differed at the nucleotide level but were identical at the protein level (PITG\_16245 = PITG\_16418; PITG\_16233 = PITG\_16240; PITG\_20934 = PITG\_20936; PITG\_16409 = PITG\_16424). The phylogenetic analysis of the complete protein sequences led to the identification of five clades. This analysis was performed in MEGA x by using the maximum likelihood method based on the JTT matrix-based model. The tree with the highest log-likelihood (-766) is shown. The bootstrapping values, which indicate the percentage of trees that had the particular branch, are shown in each branch. In the protein alignment: blue, hydrophobic residues (A, I, L, M, F, W, V, and C); red, positively charged residues (K and R); magenta, negatively charged residues (E and D); green, polar residues (N, Q, S, and T); pink, cysteine residues (C); orange, glycine residues (G); yellow, proline residues (P); cyan, aromatic residues (H and Y); and white, unconserved residues or gaps. (b) Different *Rpi-chc1* allelic variants were co-agroinfiltrated in *N. benthamiana* with a member from each PexRD12/31 clade. While variants from clade 1 recognize both PexRD12-A1 and -A2 clades, *Rpi-chc1* variants from clade 2 recognize PexRD31-B and -C. Receptors from clades 3 and 4 do not recognize any PexRD12/31 effector. A mix of *R3a* and *Avr3a* was used as a positive control.

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different PexRD12/31 members in *N. benthamiana. Rpi-chc1.1\_543-5* and *Rpi-ber1.1\_*94-2031-01 from clade 1, *Rpi-chc1.2\_543-5* and *Rpi-ber1.2\_*493-7 from clade 2, *rpi-tub1*-RH89-039-16 from clade 3, and *rpi-ber1.4\_561-2* from clade 4 were selected. As a representation from each of the clades of the PexRD12/31 effector superfamily, we selected PITG\_16245 (PexRD12-A1), PITG\_20934 (PexRD12-A2), PITG\_16235 (PexRD31-B), and PITG\_23069 (PexRD31-C). The different *Rpi-chc1* alleles were co-agroinfiltrated with the PexRD12/31 effectors in *N. benthamiana* leaves. Three days after agroinfiltration, we observed that the

members from clade 1, *Rpi-chc1.1* and *Rpi-ber1.1*, specifically recognized both PexRD12-A1 and PexRD12-A2 effectors (Figure 5b). This result showed that Rpi-chc1.1 and Rpi-ber1.1 recognize multiple members of the PexRD12 family, suggesting that Avrchc1.1 is encoded by multiple redundant paralogs. On the other hand, Rpi-chc1.2 and Rpi-ber1.2 from clade 2 specifically recognized both PexRD31-B and PexRD31-C effectors (Figure 5b). This suggests that multiple PexRD31 paralogs correspond to Avrchc1.2. The selected alleles from clades 3 and 4, rpi-tub1.3\_RH89-039-16 and *rpi-ber1.4\_561-2*, were not able to



#### Figure 6. Domain exchanges between Rpi-chc1.2 and rpi-tub1.3\_RH89-039-16.

(a) The positions of SAPs and the corresponding protein domains are indicated on top. Rpi-chc1.2 and rpi-tub1.3\_RH89-039-16 are represented as light blue and yellow bars, respectively. Below, the domain exchanges are shown. The chimeric constructs were co-agroinfiltrated with the PexRD12/31 effectors in *N. ben-thamiana* leaves. After 4 days, the HR was visible and recorded. Experiments were repeated three times with 12 inoculation sites each time. A representative leaf for the response of each chimeric construct is shown in the right panel. GUS was used as a negative control. It is concluded that the exchange of the complete LRR domain led to recognition of PexRD31. With the final construct, RH::C2\_14-19, the exchange of only nine amino acids led to the activation of the rpi-tub1.3\_RH89-036-16 protein. (b) Seven new modified receptors were derived from RH::C2\_14-19 in order to pinpoint the amino acids involved in the Rpi-chc1.2 recognition specificity. SAPs present in Rpi-chc1.2 are highlighted in blue font. Most of the SAPs are located in the solvent-exposed xxLxLxxxx motifs of LRRs 16-19. The chimeric construct is shown in the right panel. Experiments were repeated three times with 12 inoculation sites each time. GUS was used as a negative control. The modification of the tested residues of the Rpi-chc1.2 solvent-exposed specific amino acids (blue) for the corresponding amino acid present in rpi-tub1.3\_RH89-039-16 (yellow) led to the partial or complete loss of effector recognition.

recognize any of the PexRD12/31 members (Figure 5b), showing that clades 3 and 4 encode more functionally distant receptors, in agreement with the known susceptibility of RH89-039-16 to *P. infestans* (Vleeshouwers *et al.*, 2011a).

# The LRR domain of the Rpi-chc1 variants determines the PexRD12/31 effector recognition specificity

Since the allelic variants of *Rpi-chc1* could be divided into three activity groups (recognition of Avrchc1.1, Avrchc1.2, or none), while having an amino acid identity up to 96%, they provide ideal tools to investigate the Rpi-chc1 mechanism of recognition. Therefore, we performed progressive exchanges of the different receptor domains. The chimeric receptors were co-agroinfiltrated with the PexRD12/31 effectors in N. benthamiana leaves to evaluate their recognition specificity. First, we selected Rpi-chc1.2 and rpitub1.3\_RH89-039-16 as representatives of clades B and C, respectively. When aligning the protein sequences, 54 single amino acid polymorphisms (SAPs) were found and most of them were located in the LRR domain (Figure 6a). As previously mentioned, Rpi-chc1.2 specifically recognizes PexRD31-B and PexRD31-C, while rpi-tub1.3\_RH89-039-16 does not recognize any of the PexRD12/31 effectors. When the complete rpi-tub1.3 RH89-039-16 LRR domain was exchanged for the Rpi-chc1.2 LRR, the chimeric receptor RH::C2\_2-29 was able to recognize both PexRD31-B and

## Rpi-chc1 variants recognize PexRD12/31 effectors 9

PexRD31-C. Reciprocally, the exchange of the Rpi-chc1.2 LRR for the rpi-tub1.3\_RH89-039-16 in C2::RH\_2-29 led to the inability to recognize any of the PexRD12/31 effectors. This result demonstrates the importance of the LRR domain during Avrchc1.2 recognition. Additional domain exchanges were performed in order to identify the essential LRRs for effector recognition. The LRRs required for Avrchc1.2 recognition could be narrowed down with the construct RH::C2 14-19 to nine amino acid polymorphisms (Figure 6a). Due to the absence of polymorphisms in LRRs 14 and 15, we can conclude that the exchange of the amino acid polymorphisms present in LRRs 16 to 19 activates rpitub1.3 RH89-039-16 to recognize Avrchc1.2. Interestingly, the majority of these nine amino acid polymorphisms are particularly situated in the solvent-exposed domain (xxLxLxxxx) of every LRR. The exchange of the tested solvent-exposed residues led to the partial or complete loss of effector recognition, suggesting the contribution of an effector-binding surface (Figure 6b).

To understand the difference in effector recognition specificity between Rpi-chc1.1 and Rpi-chc1.2 and to explore the possibility to combine both recognitions in one receptor, we performed a similar progressive domain exchange approach between Rpi-chc1.1 and Rpi-chc1.2 (Figure 7). The exchange of the LRR domain in the chimeric receptors C1::C2\_8-29 and C2::C1\_8-29 led to a shift



Figure 7. The effector recognition specificity could be exchanged between Rpi-chc1.1 and Rpi-chc1.2.

The alignment of Rpi-chc1.1 and Rpi-chc1.2 shows that all the 41 amino acid polymorphisms (red bars) are located in the LRR domain. The chimeric constructs were co-agroinfiltrated with the PexRD12/31 effectors in *N. benthamiana* leaves. A representative leaf for the response of each chimeric construct is shown in the right panel. Experiments were repeated three times with 12 inoculation sites each time. GUS was used as a negative control. In construct C1::C2\_14-23, LRRs 16–19 appear again, determining PexRD31 recognition. The required domain exchanges of the Rpi-chc1.1 LRR are more complex and encompass almost the complete LRR.

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in effector recognition, from Avrchc1.1 to Avrchc1.2. Further exchanges revealed that LRRs 14 to 23 from Rpichc1.2 led to an opposite effector recognition pattern as the chimeric receptor C1::C2\_14-23 was only able to recognize Avrchc1.2. With reciprocal domain exchanges of Rpichc1.1 into Rpi-chc1.2, we found that LRRs 8 to 29 led to Avrchc1.1 recognition. In an attempt to further reduce the length of the exchanged sequence, the recognition of Avrchc1.1 resulted in partial (C2::C1 8-25 and C2::C1 8-23) or complete (C2::C1\_14-25 and C2::C1\_14-23) loss of recognition. Especially, when comparing the receptors C2::C1 8-29 and C2::C1\_8-25, already the modification of the last five SAPs led to the reduced recognition of Avrchc1.1. But, apparently not only the last LRRs are involved in effector recognition; also the first LRRs, from 8 to 14, are important for Avrchc1.1 recognition as C2::C1\_8-25 was able to partially recognize Avrchc1.1, while C2::C1\_14-25 did not trigger any HR. We conclude that LRRs 8 to 29 in Rpi-chc1.1 are important for Avrchc1.1 recognition, which overlap with LRRs 16-19 from Rpi-chc1.2, which are required for Avrchc1.2 recognition.

## DISCUSSION

In this study, we identified *Rpi-chc1.1* and 15 additional allelic variants from S. chacoense, S. berthaultii, S. tarijense, and S. tuberosum. Phylogenetic analysis of the encoded protein sequences revealed four clades. These four clades were not only supported by sequence similarity but also by differences in effector and P. infestans recognition. Clade 1 genes encode receptors that recognize PexRD12 effectors and includes the active orthologs Rpichc1.1, Rpi-ber1.1, and Rpi-tar1.1 (Figures 3 and 5). Clade 2 receptors could be distinguished by the recognition of the PexRD31 effectors (Figure 5). Receptors encoded by clades 3 and 4 do not recognize PexRD12/31 effectors and no other activity has been found. Interestingly, clade 3 alleles are also present in domesticated potato clones that are susceptible to late blight, e.g., RH89-039-16 (Figure 3) and the varieties Colomba and Altus (to be published elsewhere), implying that the encoded receptors are not able to effectively provide resistance against *P. infestans*.

*Rpi-ber1.1\_94-2031-01* was derived from the same accession as the previously described *Rpi-ber* (Rauscher *et al.*, 2006; Tan *et al.*, 2010; Vossen *et al.*, 2009) and *Rpi-ber1* genes (Park *et al.*, 2009). In these four studies, *Rpi-ber* and *Rpi-ber1* mapped close to marker TG63 but slightly different genetic positions were reported. The population from Park *et al.* was quite small and a single recombination event may have caused the deviating genetic distance. In the case of Tan *et al.*, a single mis-phenotyping could explain the mapping of *Rpi-ber1* are the same genes and adopt the *Rpi-ber1* naming as it is more consistent with current nomenclature for late blight resistance genes.

*Rpi-ber2*, as described by Park *et al.*, was derived from the same accession that was used in our allele-mining studies (BER493). We could not find a clade 1 *Rpi-chc1* allele from the BER493 accessions, which supports the idea that a more distantly related CNL16 member may be present that lacks sufficient match to the primer sequences, explaining the *Rpi-ber2* map position distal from TG63.

The presence of *Rpi-chc1* alleles in *S. tarijense* and *S. berthaultii* suggests a functional common ancestor existed before their speciation. However, it must be noted that the geographic locations where the accessions were found are close to each other in Bolivia. Since *S. chacoense, S. tarijense,* and *S. berthaultii* are closely related, the presence of functional *Rpi-chc1* alleles in the three species might be a result from a recent species intercrossing.

Sequence similarity among the studied allelic variants correlated with their functionality, deduced by their ability to provide late blight resistance and P. infestans effector recognition (Figures 4 and 5). This is not the first described case of R gene allelic variants across Solanum species. Rpi-blb1, Rpi-sto1, and Rpi-pta1, from the Mexican species S. bulbocastanum, Solanum stoloniferum, and Solanum papita are allelic variants that recognize the same lpiO or PexRD6 P. infestans effector (Vleeshouwers et al., 2008). Among allelic variants of late blight resistance genes (i.e., Rpi-blb3 and Rpi-hit1 that recognize Avr2 effectors), overlapping recognition specificities have been previously described (Champouret, 2010) but also more recently (Witek et al., 2021). Moreover, highly similar, but nonallelic R genes from the same CNL cluster had different recognition specificities, i.e., Rpi-vnt1, Rpi-mcg1, R9a, Ph-3 (Foster et al., 2009; Jo et al., 2015; Smilde et al., 2005; Zhang et al., 2014). In the current report, we describe for the first time that allelic variants of a late blight resistance gene show non-overlapping effector recognition specificities in the combinations that were tested. Remarkably, the recognized effectors belong to the same effector family, which is a further refinement of our insight in host-pathogen co-evolution.

When studying the Rpi-chc1 protein domain structure, we identified most of the conserved CNL motifs. Remarkably, the MADA motif (Adachi *et al.*, 2019) was not located downstream of the starting methionine, but downstream of the second methionine in position 46 of the Rpi-chc1 protein. Further research is needed to show if either or both methionines are used as translational start codons. Interestingly, we recently cloned the functional late blight resistance gene from the late blight-resistant variety Carolus (*Rpi-Carolus* gene, to be published elsewhere). Rpi-Carolus differs only at seven amino acid positions from Rpi-ber1, but its N-terminus is shorter as a stop codon is present between the first two methionine codons. This strongly suggests that translation in Rpi-Carolus starts from the second methionine while retaining biological activity.

In contrast to the relatively conserved N-termini of the proteins encoded by the *Rpi-chc1* alleles, most interallelic sequence variation localized to the LRR regions. Through domain interchange between the rpi-tub1.3\_RH89-039-16 and Rpi-chc1.2 alleles and between Rpi-chc1.1 and Rpichc1.2, we discovered that the LRR domain defines recognition specificity (Figures 6 and 7). Polymorphisms in the LRRs of some NLR receptors were previously shown to determine the effector recognition specificity (Catanzariti et al., 2010; Dodds et al., 2001; Krasileva et al., 2010; Ravensdale et al., 2012; Shen et al., 2003). In one example, a domain exchange between Rx1 and Gpa2 converted the virus resistance into nematode resistance, and vice versa (Slootweg et al., 2017). The recognition of both nematode and virus could not be combined into one chimeric receptor, as we also observed with Rpi-chc1.1 and Rpi-chc1.2. The reason for this is the overlap between the LRRs involved in recognition.

Most of the amino acids in Rpi-chc1.2 that are required for Avrchc1.2 recognition are located in the LRR solventexposed motif (xxLxLxxxx), and modification of the tested solvent-exposed amino acids led to the partial or complete loss of PexRD31 recognition (Figure 6b). The corequirement of these solvent-exposed amino acids suggests that they are involved in recognition of a particular epitope. This observation, combined with the observation of unequal distribution of SAPs, allow us to hypothesize that *Rpi-chc1* alleles evolved through insertion of a stretch of DNA into the LRR domain rather than through accumulation of independent mutations. A similar model of evolution was recently proposed for allelic variants of Rpi-amr1 (Witek et al., 2021). Such insertions may happen through unequal crossing-over with paralog sequences or through retro-transposition. Interestingly, the evolution of integrated domains in R genes has been postulated to be caused by an unknown recombination- or transposonindependent translocation mechanism (Bailey et al., 2018). The same mechanism may be active in LRR exchange to evolve recognition of non-integrated domains like guardees or decoys (Kourelis and van der Hoorn, 2018) or direct effector recognition.

Interestingly, some of the PexRD31 family members have been previously identified as one of the most rapidly diversifying and fast evolving RXLR effectors in the T30-4 genome, with  $\omega$  values higher than 1.55 (Haas *et al.*, 2009). Additionally, several members of the PexRD12/31 superfamily have recently been characterized to target the host vesicle trafficking machinery by interacting with the vesicle-associated membrane protein 72 family (Petre *et al.*, 2020). Even though both PexRD12 and PexRD31 have the same or functionally similar host targets, they are differentially expressed during *P. infestans* infection. While PexRD12 is highly expressed in all the tested isolates, PexRD31 is expressed at low levels after contact with

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potato (Figures S1 and S4). This would also explain why *Rpi-chc1.2* is not able to provide resistance against *P. infestans*, since most of the isolates have low or undetectable expression levels of *Avrchc1.2* (Figure S4). Consequently, clade A (PexRD12) may have evolved to avoid detection by *Rpi-chc1.2* while retaining its targeting of the vesicle trafficking machinery.

Another step in the co-evolution between *Rpi-chc1.1* and the PexRD12/31 family was found by analyzing the effector expression in plants expressing *Rpi-chc1.1*. The isolates that overcome the *Rpi-chc1.1* resistance no longer express PexRD12, while the expression in untransformed Desiree plants was normal and comparable to the expression of *Avrsto1* (Figure 4b). Similarly, evasion of recognition through transcriptional suppression was previously observed in plants expressing *Rpi-vnt1* infected with *P. infestans* (Pel, 2010). Once more, we confirmed the plasticity of the *P. infestans* effector secretion and the fast evolution capacity of some aggressive isolates to break down single Rpi resistances.

The introgression of single R genes drives P. infestans to evolve and evade recognition. In order to durably deploy late blight resistance in agriculture, we need novel strategies informed by knowledge of disease resistance genes in varieties, their recognition specificities, and the presence of the cognate effectors in the pathogen populations. Virulence information from the field must be rapidly translated to decision support systems (DSSs) for the risk prediction and calculation of biocide spraying intervals. Additionally, DSSs can be used to determine the R gene composition of (novel) varieties to be deployed in the next season. To meet these requirements, novel breeding strategies are needed to rapidly tailor the R gene contents of the potato varieties to the pathogen populations. In current breeding schemes, it takes 10-15 years to select a late blight-resistant potato variety. Moreover, susceptible varieties with dominant market shares will not be easy to replace. A system of varieties with flexible R gene content is needed. In other crops this has been accomplished through F1 hybrid varieties. In potato, this route has only recently been opened (Su et al., 2020) and no hybrid potato varieties have reached the market yet. A proof of principle for flexible late blight resistance varieties produced through cisgenesis was provided several years ago (Haverkort et al., 2016). Unfortunately, the EU legislation does not distinguish between cisgenic and transgenic products, making it impossible to market cisgenic varieties. In other parts of the world similar improvement technologies are being deployed to obtain late blight resistance (Ghislain et al., 2019; Habig et al., 2018). It is promising that many different national authorities now apply or consider separate oversight regulations for events that have been enriched with 'cisgenes' or 'genes with a history of safe use'. Knowledge as obtained in this study is essential to

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pursue such enrichment strategies, which can be achieved through transformation of cisgenes or though novel gene editing tools. We now know how inactive resistance genes from susceptible varieties could be repaired by replacing minimal fragments with the corresponding fragments of alleles from wild relatives. This would provide an unprecedented accuracy and speed which is not present in introgression breeding.

## **EXPERIMENTAL PROCEDURES**

#### Plant materials and growth conditions

The wild *Solanum* species used in this study are listed in Table S6 (Tan *et al.*, 2010; Vleeshouwers *et al.*, 2011a). The potato plants were maintained *in vitro* on MS20 at 24°C under a 16/8-h day/ night regime (Domazakis *et al.*, 2017). The 7650 F1 population was generated by crossing *S. chacoense* (CHC543-5)  $\times$  *S. chacoense* (CHC544-5). *Solanum tuberosum cv.* 'Desiree' was used for stable transformations of the different *Rpi-chc1.1* candidates. Four-week-old *N. benthamiana* leaves were used for agroinfiltration. The agroinfiltrated plants were kept in climate-regulated greenhouse compartments of Unifarm (Wageningen University & Research) at 20–25°C and under a 16/8-h day/night regime.

#### BAC clone isolation and sequencing

The procedure has been described in patent US9551007B2. Briefly, two different BAC libraries were produced using partial digestion of CHC543-5 genomic DNA with *Hin*dIII. Fragments larger than 100 kb were ligated into pBeloBAC or pCC1BAC arms (Epicenter). The BAC clones were collected and stored as bacterial pools of approximately 700–1000 white colonies. BAC pools were screened with selected markers and individual clones were identified using colony PCR. The ends of positive individual BACs were sequenced for the purpose of fine-mapping RH106G03T and RH137D14\_C37-7-4. The complete inserts were sequenced using shotgun sequencing of 2-kb library fragments generated by partial digestion with *Eco*RI by Macrogen Inc. (Seoul, South Korea). Assembly of the sequences resulted in contigs as indicated in Figure 1 (GenBank accession number MW383255).

# Cloning of *Rpi-chc1* allelic variants and chimeric constructs

The Rpi-chc1 allelic variants were amplified using genomic DNA from the different wild Solanum species using PCR primers as described in Table S7 and DNA polymerase with proofreading activity. The fragments were cloned into pGEM-T easy vector (Promega, Leiden, the Netherlands) for sequencing. GenBank submission numbers are provided in Table S6. The Rpi-ber1.1 and Rpitar1.1 genes were amplified using primers in the promotor and terminator. The resulting PCR fragments were cloned into pBINPLUS-PASSA (Jo et al., 2016) and were expressed in transgenic Desiree plants under the control of their native regulatory elements. For transient expression analyses, the coding sequences of the allelic variants were cloned under the Rpi-chc1.1 regulatory elements (900-bp promotor and 400-bp terminator) into pDEST using a multisite gateway protocol. Escherichia coli strain DH10ß was transformed with the gateway reaction products and clones with the correct insert were selected. Agrobacterium tumefaciens AGL1+VirG was used for transient and stable transformations of N. benthamiana leaves and S. tuberosum cv. 'Desiree'.

The chimeric constructs were cloned using the Golden Gate modular cloning principle. As acceptor vector, we used a Golden Gate-compatible version of pBINPLUS (McBride and Summerfelt, 1990), PBINPLUS-GG (van de Vossenberg *et al.*, 20192019). The final acceptor vector was constructed to contain 800-bp *Rpi-chc1.1* promoter::CDS::1000-bp *Rpi-ber* terminator (Figure S5). The different PCR fragments were amplified using the Phusion High-Fidelity PCR Kit (Thermo Scientific, Ochten, the Netherlands) and primers with *Bsa*l sites as overhang (Table S7) and purified using the DNA Clean&Concentrator Kit (Zymo Research, Freiburg im Breisgau, Germany). PCR fragments and the acceptor vector were incubated in Buffer G (Thermo Scientific) with 1 mm ATP for 30 cycles of 37°C for 5 min and 16°C for 5 min. We included an additional step at 37°C for 10 min to digest the wrongly assembled plasmids and a final step at 65°C for 20 min to heat-inactivate the *Bsa*l enzyme.

#### Hypersensitive cell death assays

Transient expression of the different receptors and PexRD12/31 effectors was performed in 4-week-old *N. benthamiana* leaves. R3a/Avr3a was used as a positive control. All the constructs were agroinfiltrated at an  $OD_{600}$  of 0.5. Each construct was agroinfiltrated twice on two leaves of four plants in at least two independent experiments. Cell death responses were observed at 3–4 days post-inoculation.

## Phylogenetic analysis of Rpi-chc1.1 homologs and the PexRD12/31 superfamily

The sequences of the PexRD12/31 effectors were retrieved from the *P. infestans* T30-4 genome (Haas *et al.*, 2009). Twenty family members were found to form the PexRD12/31 superfamily. The coding sequences of the *Rpi-chc1* variants as obtained in this study were aligned using MUSCLE and a neighbor-joining tree was constructed using MEGALIGN from the DNAstar package. The closest homolog of *Rpi-chc1.1* from the DM reference genome (Sol-tuDM10G021850.1) was used as an outgroup.

The protein sequences of PexRD12/31 effectors were aligned using Clustal OMEGA and manually edited in MEGA X (Kumar *et al.*, 2018; Sievers *et al.*, 2011). The phylogenetic relationship was inferred using the maximum likelihood method based on the JTT matrix-based model in MEGA X with 1000 bootstraps (Jones *et al.*, 1992). The tree with the highest log-likelihood is shown. The two more distant effectors PITG\_16428 and PITG\_09577 served as an outgroup.

# *Phytophthora infestans* isolates and the detached leaf assay

The P. infestans isolates used in this study (90128, IPO-C, and NL08645) were retrieved from our in-house collection. Isolates were grown at 15°C on solid rye medium in the dark (Caten and Jinks, 1968). After 2 weeks, sporulating mycelium was flooded with 20 ml of ice-cold water, adjusted to 70 zoospores/µl, and incubated at 4°C for 2-3 h. After the incubation, the detached leaves were inoculated with 10  $\mu$ l of the zoospore suspension on the abaxial side of the leaves. Detached leaves were inserted into wet floral foam. For each biological replicate the three leaflets from four leaves from two independent plants were used. Twelve spots on each leaf were inoculated with the zoospore suspension and closed in a plastic bag to maintain high humidity. The leaves were kept in a climate cell at 18°C for 5 days. Disease resistance was scored on a scale from 1 to 10 for each leaflet, where 10 = no symptoms; 9 = HR no larger than the inoculum droplet; 8 = HR lesion of up to 0.5 cm diameter; 7 = diffuse lesions up to 1 cm diameter, no sporulation, no water soaking; 5 = lesions larger than 1 cm sometimes with water soaking, no sporulation; 4 = large water-soaked lesions with sporulation only visible through binoculars; 2 = large lesions with macroscopically visible sporulation on one side of the leaflet; and 1 = large lesions with macroscopically visible sporulation on both sides of the leaflet.

#### Relative effector and R gene expression

The P. infestans effectors used in this study are listed in Table S5. The different genotypes were inoculated with the different P. infestans isolates and samples were collected after 0, 3, 8, 24, 48, 72, 96, and 120 h. Infected plant material with the different P. infestans isolates was collected and RNA was isolated using an RNA Purification Kit (Qiagen, VenIo, the Netherlands). The isolated RNA was converted into cDNA using the QuantiTect Reverse Transcription Kit (Qiagen). The primers used in this study are listed in Table S7. The expression of the different effectors in the infected material was evaluated using RT-qPCR SYBR Green (Bio-Rad, Hercules, CA, USA). The samples were heated to 95°C for 2 min, followed by 40 cycles of 15 s at 95°C, 30 s at 60°C, and 30 s at 72°C. Fluorescence was measured after each cycle. After the final amplification cycle a melting curve was calculated. Relative gene expression was calculated using the  $2^{-\Delta\Delta Ct}$  method (Livak and Schmittgen, 2001). Gene expression was normalized by dividing the relative gene expression by the relative expression of the P. infestans elongation factor 2 (ef2) gene.

#### SgRNA and CRISPR-Cas9 construct design

The CRISPOR web tool (http://crispor.org) was used to design the sgRNAs with lower off-target and higher on-target potentials (Concordet and Haeussler, 2018).

A modular cloning system based on the Golden Gate cloning technology was used to assemble the different sgRNAs and binary vectors as previously described for tomato (*Solanum lycopersicum*) mutagenesis (Engler *et al.*, 2008; Weber *et al.*, 2011). Briefly, each sgRNA was fused to the *Arabidopsis thaliana* U6-26 promoter as *AtU6-26::gRNA*. The Level 1 constructs *plCH47732-pNOS::NPTII::tOCS* and *plCH47742-p2x35S::hCas9::tNOS* and the linker *plCH41780* were used to build the Level 2 vector *plCSL4723* (Werner *et al.*, 2012). The primers used for cloning the gRNAs are listed in Table S7.

#### ACKNOWLEDGMENTS

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#### **AUTHOR CONTRIBUTIONS**

JV planned and designed the research; DML, MN, and LK performed the majority of the experiments; SK provided

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the Pex-RD set; DML and JV wrote the manuscript; RV proofread the manuscript and provided the essential research environment; SA, HS, and KS contributed by mapping, cloning, and characterizing *Rpi-chc1* allelic variants; RS, AL, and AAH contributed by identifying Avrchc1 and their differential recognition specificities by *Rpi-chc1* allelic variants.

## **CONFLICT OF INTEREST**

Some results presented in this manuscript have been included in a patent application.

#### DATA AVAILABILITY STATEMENT

Supplementary figures and tables are available as supporting information from the *TPJ* website. All described sequences have been submitted to GenBank. Accession numbers are listed in Table S6.

#### SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article.

**Figure S1**. Effector and *R* gene expression in potato leaves inoculated with *P. infestans.* 

Figure S2. Rpi-chc1.1 protein domain organization.

**Figure S3**. Localization of *PexRD12/31* effectors in the *P. infestans* T30-4 contigs.

Figure S4. PexRD12/31 effector expression in potato leaves infected with *P. infestans* from the PenSeq dataset.

Figure S5. pBINPLUS-PASSA-GG vector map.

Table S1. CRISPR-Cas9 targeting of the *Rpi-chc1.1* allele in *S. cha-coense* resistant accession CHC543-5.

 Table S2. Late blight resistance assessment of different Rpi-chc1 alleles.

 Table S3. Segregation of markers and late blight resistance of *Rpi-chc1.1* and *Rpi-chc1.2*.

 Table S4.
 Functional expression of *Rpi-chc1.1* in Desiree transgenic events correlates with responsiveness to PexRD12.

 Table S5. Phytophthora infestans effectors used in this study.

Table S6. Accession numbers of *Solanum* genotypes and *Rpi-chc1* sequences.

 Table S7. Primers used in this study.

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