# A complex resistance locus in *Solanum americanum* recognizes a conserved *Phytophthora* effector

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# 1 Abstract

Late blight caused by *Phytophthora infestans* greatly constrains potato production. Many Resistance (R) genes were cloned from wild Solanum species and/or introduced into potato cultivars by breeding. However, individual R genes have been overcome by P. infestans evolution; durable resistance remains elusive. We positionally cloned a new R gene, Rpi-amrl, from Solanum americanum, that encodes an NRC helper-dependent CC-NLR protein. Rpi-amrl confers resistance in potato to all 19 P. infestans isolates tested. Using association genomics and long-read RenSeq, we defined eight additional Rpi-amr1 alleles from different S. americanum and related species. Despite only ~90% identity between Rpi-amr1 proteins, all confer late blight resistance but differentially recognize Avramr1 orthologs and paralogs. We propose that Rpi-amr1 gene family diversity facilitates detection of diverse paralogs and alleles of the recognized effector, enabling broad-spectrum and durable resistance against P. infestans. 

# 35 Introduction

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Potato is the fourth most important directly-consumed food crop world-wide<sup>1</sup>. *Phytophthora infestans,* an oomycete pathogen, causes late blight disease in potato, and can result in complete crop failure. Disease management is primarily based on repeated fungicide applications (10-25 times per season in Europe). However, fungicideresistant races have emerged<sup>2</sup>.

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To elevate late blight resistance, *Resistance* to *Phytophthora infestans* (*Rpi*) genes were identified in wild relatives of potato and used for resistance breeding<sup>3</sup>. More than 20 *Rpi* genes have been mapped and cloned from different *Solanum* species<sup>4</sup>. All encode coiled-coil (CC), nucleotide binding (NB), leucine-rich repeat (LRR) (NLR) proteins<sup>5</sup> and some require helper NLR proteins of the NRC family<sup>6</sup>. However, most cloned *Rpi* genes have been broken by *P. infestans*<sup>7</sup>. Provision of durable late blight resistance for potato remains a major challenge.

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NLR-mediated immunity upon effector recognition activates "effector-triggered 51 52 immunity" (ETI)<sup>8</sup>. In oomycetes, all identified recognized effectors, or avirulence (Avr) 53 genes, carry a signal peptide and an RxLR motif<sup>9</sup>. 563 RxLR effectors were predicted 54 from the *P. infestans* genome, enabling identification of the recognized effectors<sup>10,11</sup>. 55 Many P. infestans effectors show signatures of selection to evade recognition by corresponding NLR proteins<sup>12</sup>. NLR genes also show extensive allelic and 56 presence/absence variation in wild plant populations<sup>13,14</sup> and known Resistance (R) 57 gene loci like Mla, L, Pi9, RPP1 and RPP13 from barley, flax, rice and Arabidopsis 58 show substantial allelic polymorphism<sup>15-18</sup>. Remarkably, different *Mla* alleles can 59 recognize sequence-unrelated effectors<sup>19,20</sup>. 60

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Technical advances like RenSeq (Resistance gene enrichment and Sequencing) and
PenSeq (Pathogen enrichment Sequencing) enable rapid definition of allelic variation
and mapping of plant *NLRs*, or discovery of variation in pathogen effectors<sup>21-23</sup>.
Combined with single-molecule real-time (SMRT) sequencing, SMRT RenSeq enabled
cloning of *Rpi-amr3* from *Solanum americanum*<sup>24</sup>. Similarly, long read and cDNA
PenSeq enabled us to identify *Avramr1* from *P. infestans*<sup>25</sup>.

69 In this study, we further explored the genetic diversity of S. americanum, and by 70 applying sequence capture technologies, we fine-mapped and cloned *Rpi-amr1* from *S*. 71 americanum, (usually) located on the short arm of chromosome 11. Multiple Rpi-amr1 72 homologs were found in different S. americanum accessions and in relatives, including 73 Solanum nigrescens and Solanum nigrum. Functional alleles show extensive allelic 74 variation and confer strong, broad-spectrum resistance to all 19 tested diverse P. 75 infestans isolates. Although differential recognition was found between different Rpi-76 amr1 and Avramr1 homologs, all Rpi-amr1 alleles recognize the Avramr1 homologs 77 from *Phytophthora parasitica* and *Phytophthora cactorum*. Our study reveals unique 78 properties of genetic variation of R genes from "non-host" species.

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# 80 **Results**

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### 82 *Rpi-amr1* maps to the short arm of chromosome 11

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84 We previously investigated S. americanum and isolated Rpi-amr3 from an accession 944750095 (SP1102)<sup>24</sup>. To discover new *Rpi-amr* genes, we characterized additional 85 14 lines of P. infestans-resistant S. americanum and close relatives S. nigrescens and 86 87 Solanum nodiflorum by crossing them to a susceptible (S) S. americanum line 88 954750186 (hereafter SP2271) (Table 1). To avoid self-pollination, a resistant parent 89 was always used as a pollen donor. All the corresponding F1 plants (6-10 per cross) 90 were resistant in a detached leaf assay (DLA) (Table 1). Around 60-100 F2 progeny 91 derived from each self-pollinated F1 plant were phenotyped by DLA using P. infestans 92 isolate 88069<sup>26</sup>. The F2 progenies that derived from the resistant parents with working 93 numbers SP1032, SP1034, SP1123, SP2272, SP2273, SP2360, SP3399, SP3400, 94 SP3406, SP3408 and SP3409 segregated in a ratio suggesting the presence of a single 95 (semi-) dominant resistance gene (fitting 3:1 or 2:1 [likely due to segregation distortion], 96 R:S - resistant to susceptible - ratio). Two crosses showed a 15:1 segregation (resistant 97 parent SP2300 and SP2307), suggesting the presence of two unlinked resistance genes, 98 while SP1101 showed no susceptible plants in 100 individuals, suggesting the presence 99 of three or more resistance genes.

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# 101 Table 1. S. americanum, S. nodiflorum and S. nigrescens accessions used in this

102 study and the corresponding *Rpi-amr1* homologs

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#### Table 1

Accession	Working name	Species	Reported origin	Source	Late blight resistance	<i>Rpi-amr1</i> homolog	Similarity	Cloning method
954750186	SP2271	S. americanum	Brazil	RU	Susceptible			
954750184	SP2273	S. americanum var. patulum	unknown	RU	Resistant	Rpi-amr1-2273	100%	Map-based cloning
sn27	SP1032	S. americanum sensu lato	China	BGS	Resistant	Rpi-amr1-1032	92.8%	Association genomics
Veg422	SP1034	S. americanum sensu lato	unknown	NN	Resistant	Rpi-arm1-2273	100%	Association genomics
A54750014	SP1101	S. americanum sensu lato	unknown	RU	Resistant	Rpi-amr1-1101	89.4%	SMRT RenSeq
A14750006	SP1123	S. americanum sensu lato	unknown	RU	Resistant	Rpi-amr1-1123	91.8%	Association genomics
954750174	SP2272	S. americanum	unknown	RU	Resistant	Rpi-amr1-2272	89.4%	Association genomics
SOLA 226	SP2300	S. americanum	Cuba	IPK	Resistant	Rpi-amr1-2300	90.4%	SMRT RenSeq
SOLA 425	SP2307	S. americanum	America	IPK	Resistant	Rpi-amr1-2307	91.7%	Association genomics
Wang 2059	SP2360	S. americanum	China	NHM	Resistant	Rpi-arm1-2273	100%	Association genomics
A14750138	SP3399	S. americanum	unknown	RU	Resistant	Rpi-amr1-2272	89.4%	Association genomics
A14750130	SP3400	S. nodiflorum	unknown	RU	Resistant	Rpi-amr1-2273	100%	Association genomics
944750261	SP3406	S. nigrescens	Bolivia	RU	Resistant	Rpi-amr1-3406	92.5%	Association genomics
954750172	SP3408	S. nigrescens	Bolivia	RU	Resistant	Rpi-amr1-3408	92.6%	Association genomics
A14750423	SP3409	S. nigrescens	Mauritius	RU	Resistant	Rpi-amr1-3409	89.5%	SMRT RenSeq

RU - Radboud University, Nijmegen, The Netherlands

IPK - IPK Gatersleben, Germany

NHM - Natural History Museum, London, United Kingdom

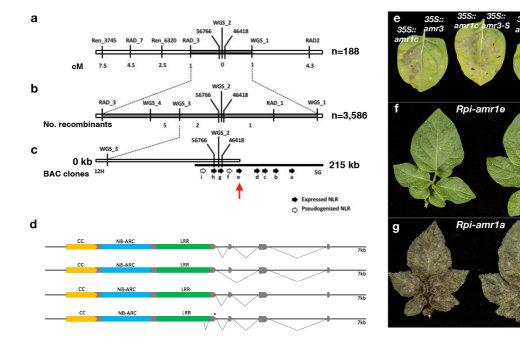
BGS - Shanghai Botanical Garden, Shanghai, China

104 NN - Nicky's Nursery Ltd, Kent, United Kingdom

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106	To identify Rpi genes from these resistant S. americanum accessions, we prioritized an
107	F2 population derived from resistant parent SP2273 and named the corresponding gene
108	Rpi-amr1. Using markers from RenSeq, genotyping by sequencing (RAD markers) and
109	Whole Genome Shotgun sequencing (WGS), the Rpi-amr1 gene was mapped in a small
110	population (n=188 gametes) to the short arm of chromosome 11, between markers
111	RAD_3 and WGS_1 (Fig. 1a, Table S1, S2). We expanded the mapping population and
112	developed a PCR marker WGS_2 that co-segregated with resistance in 3,586 gametes
113	(Fig. 1b, Table S2). To generate the physical map of the target interval from SP2273, a
114	BAC library was generated. Two BAC clones (12H and 5G) covering the target interval
115	were isolated and sequenced on the PacBio RSII platform, and assembled into a single
116	contig of 212 kb (Fig. 1c). We predicted 11 potential coding sequences on the BAC_5G,
117	nine of which encode NLR genes (Fig. 1c). These NLR genes belong to the CNL class
118	and have 80-96% between-paralog identity.

- 119 To define which of these *NLR* genes are expressed, cDNA RenSeq data of the resistant
- 120 parent SP2273 were generated and mapped to the BAC 5G sequence. Seven out of nine
- 121 NLR genes were expressed. These genes Rpi-amr1a, b, c, d, e, g and h were tested
- 122 as candidate genes for *Rpi-amr1* (Fig. 1c).
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#### 126 Fig. 1. Map-based cloning of *Rpi-amr1* and its resistance to *P. infestans*.

- (a) Mapping of *Rpi-amr1* in a small F2 population (n=188 gametes); the names of the markers and genetic distances are shown above or below the bar.
- (b) Fine mapping of *Rpi-amr1* in the F2 population of 3,586 gametes. The names of the markers and the number of recombinants are shown above or below the bar.
- (c) Physical map of the target *Rpi-amr1* interval based on the assembled BAC contig. The markers present on the BAC are shown. The predicted NLR genes are depicted as black arrows (expressed NLRs) or empty arrows (pseudogenized NLRs). *Rpi-amr1* (formerly *Rpi-amr1e*) is indicated by a red arrow.
- (d) Four *Rpi-amr1* transcripts detected by 3' RACE PCR.
- (e) Leaves of *N. benthamiana* plants were infiltrated with the binary vector pICSLUS0003::35S overexpressing either the late blight resistance gene *Rpi-amr3* (positive control), one of seven *Rpi-amr1* candidates, or the non-functional *Rpi-amr3-S* (negative control). Leaves were inoculated with *P. infestans* strain 88069 24 h after infiltration. Only leaves infiltrated with *Rpi-amr3 and Rpi-amr1e* (pictured) showed reduced pathogen growth, whereas *P. infestans* grew well in the presence of the remaining *Rpi-amr1* candidates. Only *Rpi-amr1c* is shown as the phenotype of all other non-functional candidate genes was indistinguishable. Photographs were taken 9 dpi.
- (f) Transgenic potato cv. Maris Piper which expresses *Rpi-amr1* under the native regulatory elements is resistant to *P. infestans* isolate 88069 (top), displaying no symptoms at the spot of inoculation. Each leaflet was inoculated with a droplet containing approximately 1,000 zoospores; photographs were taken 9 dpi.
- (g) The control plants carrying the non-functional candidate *Rpi-amr1a* show large necrotic lesions and sporulation. Each leaflet was inoculated with a droplet containing approximately 1,000 zoospores; photographs were taken 9 dpi.

# 152 *Rpi-amr1e* confers resistance in *Nicotiana benthamiana* and cultivated potato

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154 To test the function of the seven candidate genes, we cloned their open reading frames 155 into a binary expression vector under control of the 35S promoter. Rpi-amr3 was used 156 as a positive control and the non-functional *Rpi-amr3-S* was used as a negative control. 157 The constructs carrying each of the seven candidate genes were transiently expressed after Agrobacterium infiltration into N. benthamiana leaves, which were subsequently 158 inoculated with the P. infestans isolate 88069 as described<sup>24</sup>. P. infestans growth was 159 160 observed six days post inoculation (dpi). Only 35S::Rpi-amr1e-infiltrated leaves 161 showed reduced pathogen growth at 6 dpi compared to other candidate genes like *Rpi*amrlc, or negative control *Rpi-amr3-S*. (Fig. 1e). Hence, we conclude that *Rpi-amrle* 162 163 is the functional *Rpi-amr1* (hereafter) gene from *S. americanum* SP2273. 164

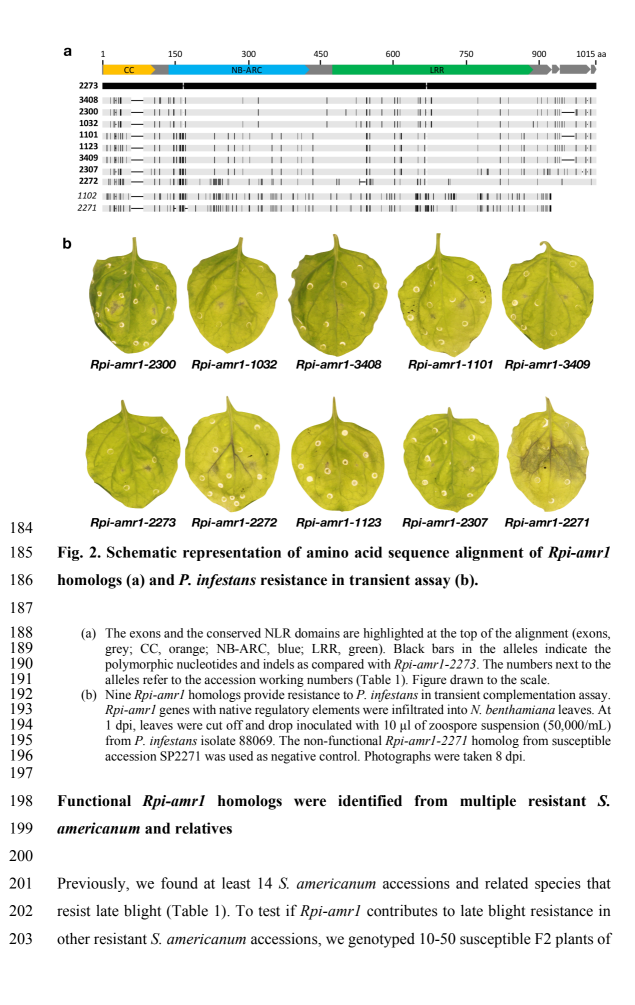
To test if *Rpi-amr1* confers late blight resistance in potato, we cloned it with its native promoter and terminator, and generated transgenic potato cultivar Maris Piper plants carrying *Rpi-amr1*. A non-functional paralog *Rpi-amr1a* was also transformed into Maris Piper as a negative control. As in the transient assay, stably transformed *Rpiamr1* lines resisted *P. infestans* 88069 in potato (Fig. 1f), but *Rpi-amr1a*-transformed plants did not (Fig. 1g).

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#### 172 *Rpi-amr1* is a four exon CC-NLR

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174 To characterize the structure of *Rpi-amr1*, we mapped the cDNA RenSeq data to the full length Rpi-amr1 gene, and found four alternatively spliced forms of Rpi-amr1. The 175 176 most abundant form, supported by >80% of reads, comprises four exons encoding a 177 protein of 1,013 amino acids. This was confirmed with 3' RACE PCR (Fig. 1d). The 178 Rpi-amr1 is a typical CC-NB-LRR resistance protein, with a coiled-coil domain (CC; 179 amino acids 2-146), nucleotide binding domain (NB-ARC; amino acids 179-457) and 180 leucine-rich repeats (LRR; located between amino acids 504-900) which are all positioned in the first exon (1-918 aa, Fig. 2a). The remaining three short exons (amino 181 acids 919-943, 944-1002 and 1,003-1,013) lack homology to any known domains. No 182 integrated domains<sup>27</sup> were found in the Rpi-amr1 protein. 183



204 the populations derived from resistant accessions, with *Rpi-amr1* linked markers 205 (markers 3745 and 56766, Fig. 1 and Table S2). We found that in SP1032, SP1034, 206 SP1123, SP2272, SP2307, SP2360, SP3399, SP3400, SP3406 and SP3408, resistance 207 is linked to the *Rpi-amr1* locus. To test if in these accessions the resistance is conferred 208 by functional *Rpi-amr1* homologs, we performed SMRT RenSeq-based *de novo* 209 assembly of each resistant accession, and looked for homologs with the greatest identity 210 to Rpi-amr1. For accessions SP2307, SP3399 and SP3406, we also used cDNA RenSeq 211 to monitor their expression. We mapped *de novo* contigs to the coding sequence of *Rpi*-212 *amr1* allowing for 15% mismatches and gaps, and selected the closest homolog as a 213 candidate *Rpi-amr1* ortholog (Table S3). In three resistant parents, namely SP1034, 214 SP2360 and SP3400, the functional alleles showed 100% identity at the amino acid 215 level to Rpi-amr1, while amino acid sequences from the remaining accessions had as 216 little as 89% identity to the functional Rpi-amr1 (Table S3). As described previously, 217 we transiently expressed the closest related candidate *Rpi-amr1* homologs in *N*. 218 benthamiana leaves followed by DLA with P. infestans isolate 88069, and verified their 219 functionality. The unique homologs of *Rpi-amr1-2273* were named as *Rpi-amr1-1032*. 220 *Rpi-amr1-1123*, *Rpi-amr1-2272*, *Rpi-amr1-2307* and *Rpi-amr1-3408*.

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For some accessions, like SP1101 and SP2300, the *Rpi-amr1*-linked markers gave ambiguous results, so we directly performed bulked segregant analysis (BSA) and RenSeq. Additional *Rpi-amr1* co-segregating paralogs, *Rpi-amr1-1101* and *Rpi-amr1-2300*, were identified and verified in transient assays as above (Fig. 2b).

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227 Similarly, we inspected an F2 population derived from S. nigrescens accession SP3409 228 (Table 1). We applied BSA RenSeq and SMRT RenSeq to the resistant parents and F2 229 segregating population, and we found five candidate NLRs belonging to the same Rpi-230 amr1 clade, all of which are expressed. The five candidates were cloned, and transient 231 assays verified one of them as a functional Rpi-amr1 homolog, Rpi-amr1-3409. 232 However, *Rpi-amr1-3409* does not co-segregate with *Rpi-amr1-*linked markers. We 233 used GenSeq sequence capture-based genotyping (Chen et al. 2018), and found that 234 *Rpi-amr1-3409* locates on chromosome 1, based on the potato DM reference genome<sup>28</sup>. 235 This result suggests that a fragment of DNA that locates on distal end of the short arm 236 of chromosome 11 in other resistant accessions was translocated to the distal end of the 237 long arm of chromosome 1 in SP3409.

238 When the full-length amino acid sequences of nine *Rpi-amr1* homologs were aligned,

the polymorphisms between different functional alleles were found to be distributed

through all domains including the LRR region (Fig. 2a and Fig. S1).

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Taken together, by using BSA RenSeq, SMRT RenSeq, cDNA RenSeq, association
genomics and GenSeq, we cloned eight additional functional *Rpi-amr1* homologs from
different resistant accessions, of which all confer resistance to *P. infestans* 88069 in
transient assays. The closest *Rpi-amr1* homolog from susceptible parent SP2271 does
not confer resistance (Fig. 2b).

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#### 248 *Rpi-amr1* is present in hexaploid *S. nigrum* accessions

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250 Most S. nigrum accessions are highly resistant to P. infestans and S. nigrum has been reported to be a "non-host" to P. infestans<sup>29</sup>, even though rare accessions are 251 252 susceptible<sup>30</sup>. S. americanum may be the diploid ancestor of hexaploid S. nigrum<sup>31</sup>. To 253 test if *Rpi-amr1* also contributes to late blight resistance in *S. nigrum*, we amplified and 254 sequenced the first exon of *Rpi-amr1* from four resistant and one reported susceptible S. nigrum accessions<sup>30</sup>. From three resistant accessions (SP1095, SP1088 and SP1097; 255 256 Table S4), we amplified sequences with >99% nucleotide identity to S. americanum 257 Rpi-amr1-2273 (Fig. S2). Rpi-amr1-1104 was more polymorphic, with 96.7% 258 nucleotide identity to Rpi-amr1-2273, and primers used for allele mining did not 259 amplify anything from the susceptible line SP999. These data suggest that *Rpi-amr1* 260 homologs are present in some S. nigrum accessions and were most likely inherited from 261 S. americanum.

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# 263 *Rpi-amr1* confers broad-spectrum late blight resistance in cultivated potato

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To test the scope of late blight resistance conferred by *Rpi-amr1* and its homologs, we generated stably transformed transgenic potato cv Maris Piper plants carrying *Rpiamr1-2272* and *Rpi-amr1-2273*, the most diverged of the homologs (Table S3), and inoculated them by DLA with 19 *P. infestans* isolates from UK, the Netherlands, Belgium, USA, Ecuador, Mexico and Korea (Table 2). Many of the tested *P. infestans* isolates can defeat multiple *Rpi* genes (Table 2). Our DLAs show that Maris Piper carrying *Rpi-amr1-2272* or *Rpi-amr1-2273* resist all 19 tested *P. infestans* isolates,

- 272 while the wild-type Maris Piper control is susceptible to all of them. This indicates that
- 273 Rpi-amr1 confers broad-spectrum resistance against diverse P. infestans races.
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#### 275 Table 2. Phenotypes of potato plants stably transformed with Rpi-amr1-2272 and

- 276 *Rpi-amr1-2273* after inoculation with multiple isolates of *P. infestans*.
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Isolate	Rpi-amr1- 2272	Rpi-amr1- 2273	Maris Piper	Origin	Race <sup>e</sup>
NL00228	R	R	S	The Netherlands	1.2.4.7
US23	R	R	S	USA	n.a.
3928A <sup>a</sup>	R	R	S	UK	$1.2.3.4.5.6.7.10.11^{\rm f}$
EC3626 <sup>b</sup>	R	R	S	Ecuador	n.a.
NL14538°	R	R	S	The Netherlands	n.a.
NR47UH <sup>d</sup>	R	R	S	UK	$1.3.4.7.10.11^{\rm f}$
T30-4	R	R	S	The Netherlands	n.a.
USA618	R	R	S	USA	1.2.3.6.7.10.11
KPI15-10	R	R	S	Korea	n.a.
IPO-C	R	R	S	Belgium	1.2.3.4.5.6.7.10.11
PIC99189	R	R	S	Mexico	1.2.5.7.10.11
UK7824	R	R	S	UK	n.a.
PIC99177	R	R	S	Mexico	1.2.3.4.7.9.11
VK98014	R	R	S	The Netherlands	1.2.4.11
NL08645	R	R	S	The Netherlands	n.a.
PIC99183	R	R	S	Mexico	1.2.3.4.5.7.8.10.11
NL11179	R	R	S	The Netherlands	n.a.
EC1 <sup>b</sup>	R	R	S	Ecuador	1.3.4.7.10.11
NL01096	R	R	S	The Netherlands	1.3.4.7.8.10.11

278 279 <sup>a</sup> Clonal lineage EU\_13\_A2, or "Blue13"

<sup>b</sup> Overcomes *Rpi-vnt1* 

280 <sup>c</sup> Overcomes *Rpi-vnt1* and partially *Rpi-blb1*, *Rpi-blb2* 

281 <sup>d</sup>Clonal lineage EU 6 A1, commonly known as "Pink6"

282 e Summarized in<sup>32</sup>

283 <sup>f</sup>See<sup>33</sup>

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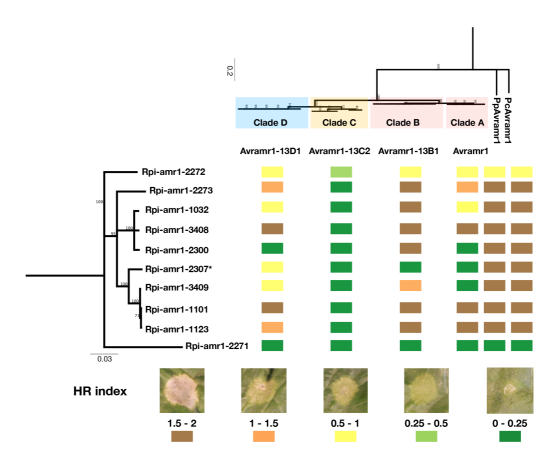
#### 286 Differential recognition by *Rpi-amr1* alleles of *Avramr1* homologs

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288 Avramr1 (PITG 07569) was identified in P. infestans race T30-4 by long-read and 289 cDNA PenSeq, and multiple Avramr1 homologs were identified in four P. infestans isolates and classified into four subclades<sup>25</sup>. To investigate if all nine cloned *Rpi-amr1* 290 291 homologs could recognize diverse Avramr1 homologs from different P. infestans 292 isolates, in addition to Avramr1 from race T30-4 that corresponds to clade A, we 293 synthesized three Avramr1 homologs Avramr1-13B1, Avramr1-13C2 and Avramr1-294 13D1 from isolate 3928A (EU 13 A2, commonly known as "Blue 13"), corresponding 295 to clades B, C and D, respectively (Fig. 3). We also synthesized the Avramr1 homologs from *P. parasitica* and *P. cactorum*<sup>25</sup>. These six *Avramr1* homologs were co-expressed 296 297 in *N. benthamiana* by agro-infiltration in all possible combinations with nine functional 298 *Rpi-amr1* homologs and the non-functional *Rpi-amr1-2271* as a negative control (Fig. 299 3).

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301 We found that different combinations of *Rpi-amr1* alleles and *Avramr1* homologs led 302 either to strong, weak or no HR phenotype in transient assay, but the non-functional 303 Rpi-amr1-2271 allele failed to recognize any Avramr1 homologs (Fig. 3). Rpi-amr1-304 2300 and Rpi-amr1-2307 recognized one Avramr1 homolog each, but others detected 305 Avramr1 homologs from more than one clade. Clade C, represented here by Avramr1-306 13C2, is usually not expressed<sup>25</sup>, and when expressed from 35S promoter, this effector 307 was not recognized by most *Rpi-amr1* homologs, though a weak HR was observed upon 308 co-expression with Rpi-amr1-2272. Avramr1-13D1 belongs to Clade D, which is absent in T30-4 but present in four other sequenced isolates<sup>25</sup>, and was recognized by 309 310 all but one (Rpi-amr1-2300) homologs in the transient assay. Surprisingly, two 311 Avramr1 homologs from P. parasitica and P. cactorum are strongly recognized by all 312 functional Rpi-amr1 homologs, apart from Rpi-amr1-2272 which showed a weaker HR 313 (Fig. 3).



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#### 315 Fig. 3. Differential recognition of *Rpi-amr1* and *Avramr1* homologs.

Four *Avramr1* homologs representing clades A-D, and *P. parasitica* and *P. cactorum* homologs were coinfiltrated with ten Rpi-amr1 homologs, including a non-functional homolog *Rpi-amr1-2271*, into *N. benthamiana* leaves. Colours from green to brown represent the strength of HR scored from 0 to 2 (see
bottom panel). N=3.

Left: phylogenetic tree of nine functional *Rpi-amr1* homologs and non-functional homolog *Rpi-amr1-*2271. Top: phylogenetic tree of *Avramr1* homologs from four isolates of *P. infestans*.

322 \* Stable *Rpi-amr1-2307 N. benthamiana* transformants show HR upon transient expression of *Avramr1* 323 and *Avramr1-13B*1.

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325 Collectively, our data shows that *Rpi-amr1/Avramr1* homolog pairs provoke 326 quantitatively and qualitatively different HRs, but all functional *Rpi-amr1* homologs

- detect at least one *Avramr1* homolog from *P. infestans* isolate 3928A.
- 328

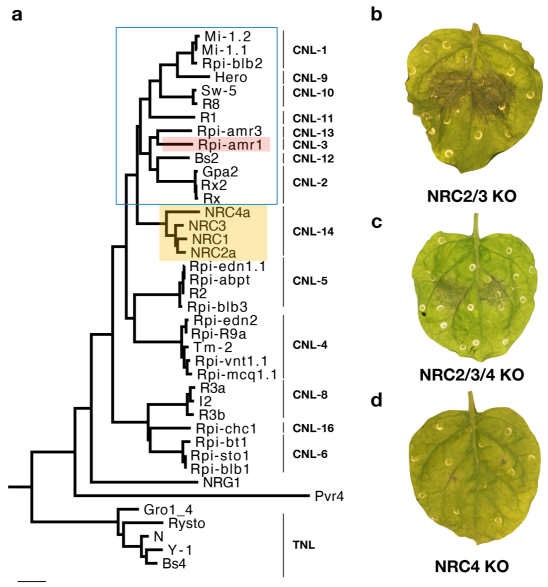
#### 329 Both *Rpi-amr1*-mediated resistance and effector recognition are NRC2 or NRC3

- 330 dependent
- 331
- 332 We generated a phylogenetic tree for representative Solanaceae NLR proteins. Rpi-
- amr1 is grouped with clade CNL-3, from which no functional resistance genes were
- 334 previously cloned (Fig. 4a). This phylogenetic affiliation suggested that Rpi-amr1 is

335 likely to depend on the helper NRC clade because CNL-3 is among the large super-336 clade of NRC-dependent sensors (Fig. 4a)<sup>6</sup>.

337 To test this hypothesis, we transiently expressed *Rpi-amr1-2273* together with *PpAvramr1* in NRC4, NRC2/3 or NRC2/3/4 knock out *N. benthamiana* leaves<sup>34,35</sup> (Fig. 338 339 S3). The HR phenotype was abolished in NRC2/3 and NRC2/3/4 knockout plants (Fig. 340 S4 c and b), but not in NRC4 knock-out or wild-type plants (Fig. S4 d and a). The HR 341 was recovered when NRC2 or NRC3 was co-expressed in the NRC2/3/4 or NRC2/3 342 knock out plants, but co-expression of NRC4 did not complement the loss of HR 343 phenotype in NRC2/3/4 knockout plants. (Fig. S4 b and c). We further showed that also 344 Rpi-amr1 mediated resistance is dependent on NRC2 or NRC3 but not NRC4, as 345 transient expression of Rpi-amr1-2273 followed by P. infestans infection restricted 346 pathogen growth only in NRC4 knockout N. benthamiana plants (Fig. 4b). These data 347 indicate that both the effector recognition and resistance conferred by *Rpi-amr1* is 348 NRC2 or NRC3 dependent.

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#### 351 Fig. 4. Rpi-amr1 is NRC2 or NRC3 dependent.

- (a) Phylogenetic analysis of Rpi-amr1 protein and other functional Solanaceae NLR proteins. The NLR clades shown here are as described previously<sup>24</sup>, the NRC-dependent sensor clades are marked by blue box.
- (b) Transient expression of *Rpi-amr1-2273* in *NRC2/NRC3* double knockout *N. benthamiana*, followed by zoospore inoculation of *P. infestans* isolate 88069, results in large necrotic lesions indicating lack of resistance.
  - (c) Transient expression of *Rpi-amr1-2273* in *NRC2/NRC3/NRC4* triple knockout *N. benthamiana*, followed by zoospore inoculation of *P. infestans* isolate 88069, results in large necrotic lesions indicating lack of the resistance.
- (d) Transient expression of *Rpi-amr1-2273* in *NRC4* knockout *N. benthamiana*, followed by zoospore inoculation of *P. infestans* isolate 88069 results in small necrotic lesions indicating resistance.
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- 365

# 366 High allelic diversity at *Rpi-amr1* was generated through inter-paralog and 367 ortholog sequence exchange

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369 *Rpi-amr1* alleles show relatively high nucleotide diversity ( $\pi$ =0.04), which could be an 370 indication of balancing or diversifying selection (Table S5). In addition, Rpi-amr1 371 alleles differ in their recognition of the Avramr1 homologs (Fig. 3) which is also 372 consistent with selection in a host-parasite co-evolutionary arms race. To test the 373 hypothesis that allelic polymorphism at *Rpi-amr1* results from diversifying selection, 374 we calculated diversity statistics and performed a McDonald-Kreitman test on both Rpi-375 amrl alleles and Avramrl homologs. As expected, Avramrl homologs show a signature 376 consistent with balancing selection (Tajima's D = 2.27) (Table S5). Remarkably, 377 despite the high nucleotide diversity, no clear signals of balancing or diversifying 378 selection were detected for *Rpi-amr1* (Tajima's D = 0.09083) (Table S5). Aligning the 379 *Rpi-amr1* alleles against the reference and scrutinizing the sequences in more detail 380 provided further insights. The nucleotide similarity of alleles varies markedly across 381 the *Rpi-amr1* homologs (Fig. 2a and Table S3); this pattern is consistent with occasional 382 recombination between highly diverged alleles or paralogs.

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384 To test whether recombination could explain the observed polymorphisms in *Rpi-amr1* 385 alleles, we predicted the possible recombination events using 3SEQ. Several 386 recombination events were detected between *Rpi-amr1* orthologs from different S. 387 americanum accessions, and Rpi-amr1 paralogs from SP2273 (Table S6). Some 388 sequence exchanges were visualized using HybridCheck (Fig. S5)<sup>36</sup>, and these data 389 suggest that sequence exchange occurred between functional Rpi-amr1 alleles and 390 paralogs. To confirm these findings, we mapped all cloned *Rpi-amr1* CDS back to the 391 BAC 5G sequence from accession SP2273 (Fig. S6). As expected, some Rpi-amr1 392 homologs (e.g. SP2300 and SP2272) show a perfect match with the fourth NLR, and 393 show a distribution of high identity that reflects the intron-exon structure. For some 394 homologs (e.g. 2271), 5' end sequences match different NLR sequences on the 395 BAC 5G and for others (e.g. 2275) part of the sequence is highly diverged from 396 BAC 5G. Taken together, our results indicate that the polymorphism of *Rpi-amr1* 397 alleles appears to have arisen partly due to sequence exchange between highly diverged 398 alleles and paralogs, and not just through mutation accumulation.

# 399 **Discussion**

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401 Achieving durable resistance is the ultimate goal of resistance breeding. Here, we report 402 significant progress towards durable resistance against potato late blight. Most cloned 403 late blight resistance genes derive from wild potatoes, and many have been overcome by one or more *P. infestans* strains<sup>37</sup>. Conceivably, resistance to *P. infestans* in nearly 404 405 all S. americanum and S. nigrum accessions is due to multiple NLR genes, as zoospores 406 from *P. infestans* can germinate on *S. nigrum* leaves but penetration is stopped by strong 407 HR<sup>29,38</sup>. *Rpi* genes from plant species that only rarely support pathogen growth have 408 likely not participated, or are no longer participating, in an evolutionary arms race with 409 P. infestans, and hence, the pathogen's effectors have not (yet) evolved to evade 410 detection by these *Rpi* genes. Under this scenario, a pre-existing standing variation in 411 the pathogen for overcoming such *Rpi* genes is either absent or extremely rare. This 412 makes such genes promising candidates for provision of broad-spectrum and durable 413 late blight resistance, provided they are not deployed alone which facilitates one-step 414 genetic changes in the pathogen to evade them, but rather in combination with other 415 genes, as in the source  $plant^{39}$ .

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417 We report here a novel, broad-spectrum S. americanum resistance gene, Rpi-amr1. We 418 also identified eight additional Rpi-amr1 alleles from different S. americanum 419 accessions and relatives, including one *Rpi-amr1* allele that translocated to the long arm 420 of chromosome 1. Allele mining also suggested the presence of *Rpi-amr1* homologs in 421 S. nigrum. All nine cloned *Rpi-amr1* alleles confer late blight resistance in transient 422 assays in N. benthamiana, and both Rpi-amr1-2272 and Rpi-amr1-2273 in potato cv 423 Maris Piper background confer resistance to all 19 tested P. infestans isolates from 424 different countries, many of which overcome other *Rpi* genes. Thus, *Rpi-amr1* is widely 425 distributed in germplasm of S. americanum, its relatives and S. nigrum, and may 426 contribute to the resistance of nearly all accessions to P. infestans.

427

Many plant *R* genes and their corresponding *Avr* genes evolved differential recognition
specificities with extensive allelic series for both *R* gene and *Avr* genes. Examples
include *ATR1* and *RPP1* or *ATR13* and *RPP13* from *Hyaloperonospora arabidopsidis*

431 and Arabidopsis<sup>9</sup>, Avr567 and L genes from the rust Melampsora lini and flax<sup>40</sup>, and

432 multiple and diverse recognized effectors from barley powdery mildew and *Mla* from 433 barley. In the same manner, *Avramr1* and its homologs from several *P. infestans* races<sup>25</sup> 434 were found to be differentially recognized by high allelic variation at the *Rpi-amr1* gene. 435 Remarkably though, the nucleotide diversity of the *R* gene did not show any of the 436 hallmarks of diversifying or balancing selection.

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438 Rather than through mutation accumulation, the high allelic variation observed at *Rpi*-439 *amr1* appears to have been generated partly by recombination between significantly 440 diverged alleles and paralogs. The recombination events are likely to be rare relative to 441 the mutation rate, given that the alleles carry many polymorphisms. This evolutionary 442 scenario can explain the observed mosaic-like structure of high and low sequence similarities when the Rpi-amr1 alleles were mapped against the contig based on two 443 444 overlapping BAC clones. The deep coalescence of alleles that is implicit in this scenario 445 can be generated by balancing selection, but we did not find evidence of such selection 446 when analysing the nucleotide substitution patterns. Recombination between *Rpi-amr1* 447 alleles could have eroded this signature of selection, as has been observed also in Rp1 resistance genes in grasses<sup>41</sup> and in the vertebrate immune genes of the major 448 histocompatibility complex (MHC)<sup>42,43</sup>. Nucleotide sequence diversity across the Rpi-449 450 amr1 alleles is correlated with only slight differences in Avramr1 recognition 451 specificity. Rpi-amr1 alleles can even recognize multiple Avramr1 paralogs from a 452 single *P. infestans* strain, a scenario that might elevate durability of resistance. Since 453 the S. americanum population recognizes multiple Avramr1 alleles and paralogs, small 454 mutational changes in Avramr1 gene are unlikely to suffice to escape detection, which 455 makes resistance-breaking less likely, thus promoting evolutionary durability of *Rpi*-456 amr1. We hypothesise that this enhanced recognition capacity could be key to the 457 evolution of "non-host" resistance, offering an escape from the coevolutionary arms 458 race. Conceivably, stacking Rpi-amr1 alleles in cis could extend the recognition 459 specificities, which could potentially lead to even more durable late blight resistance.

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Intriguingly, two *Avramr1* homologs from *P. parasitica* and *P. cactorum* are recognized by all *Rpi-amr1* homologs. Presumably, these genes have been under even less selection pressure to evade *Rpi-amr1* recognition. This result indicates that *Rpi-amr1* has the potential to provide "non-host" type resistance in *S. americanum* against multiple oomycete pathogens like *P. parasitica* and *P. cactorum*, which can infect a

wide range of hosts. As both the resistance and effector recognition of *Rpi-amr1* are *NRC2* or *NRC3* dependent, co-expression of *NRC2* or *NRC3* with *Rpi-amr1* might

468 enable it to confer resistance to other *Phytophthora* species outside the *Solanaceae*.

469

470 In summary, we cloned *Rpi-amr1*, a broad-spectrum *Rpi* gene that contributes to the 471 strong resistance of nearly all S. americanum accessions to late blight. The apparent 472 redundancy across the *Rpi-amr1* gene family may serve an evolutionary function by 473 broadening the scope for recognizing multiple Avramr1 alleles and paralogs, and potentially reducing the probability of evolution of resistance-breaking strains. 474 475 Stacking this type of *Rpi* gene with additional *Rpi* genes might help to turn host plants such as potato into non-hosts for late blight, enabling broad-spectrum and durable 476 477 resistance.

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# 482 Methods

483 Methods and associated references are in supplementary information.

484

# 485 Accession codes

486 Supporting raw reads and annotated BAC sequences were deposited in European Nucleotide

- 487 Archive (ENA) under project number PRJEB38240.
- 488

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609

# 610 Author contributions:

- 611 K.W., X.L., F.J., R.S., C.O. and J.D.G.J. designed the study. K.W., X.L., H.S.K., F.J., A.I.W.,
- 612 S.B., W.B., L.T. and T.S., performed the experiments. K.W., X.L., H.S.K., F.J., A.I.W., B.S.,
- R.S., C.O., S.F., and J.M.C. analysed the data. K.W., X.L., H.S.K., F.J. and J.D.G.J. wrote the
- 614 manuscript with input from all authors. V.G.A.A.V., B.B.H.W, C.-H.W., H.A. and S.K.

615 contributed resources. K.W., X.L and H.S.K. made equivalent contributions and should be 616 considered joint first authors. All authors approved the manuscript.

617

# 618 **Conflict of interest:**

K.W., H.S.K., F.G.J. and J.D.G.J. are named inventors on a patent application
(PCT/US2017/066691) pertaining to *Rpi-amr1* that was filed by the 2Blades Foundation on
behalf of the Sainsbury Laboratory.

622

# 623 Supplementary files:

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625 Fig. S1: Alignment of Rpi-amr1 proteins, including non-functional homolog from SP2271.

626 Fig. S2: Alignment of *Rpi-amr1-2273* and *Rpi-amr1* DNA sequences from *S. nigrum*.

627 Fig. S3: Genotypes and phenotypes of *NRC2/3* knockout *N. benthamiana*. (a). Amplicon

628 sequencing results of the *NRC* loci of the *NRC2/3* knockout *N. benthamiana* line *nrc23*\_1.3.1.

629 Sequences of the sgRNAs are marked in red; (b). The *NRC2/3* knockout line (*nrc23*\_1.3.1) did

not exhibit any growth defects when compared to the wild type plants. Six-week-old wild type

631 and *NRC2/3* knockout *N. benthamiana* lines were used in the photograph.

632 Fig. S4: The effector recognition of *Rpi-amr1* is NRC2 or NRC3 dependent. The *Rpi-amr1* 

and *Pp-Avramr1* were co-expressed by agro-infiltration on (a) wild type *N. benthamiana*; (b)

634 NRC2/3/4 knockout line; (c) NRC2/3 knockout line and (d) NRC4 knockout line. *NRC2*, *NRC3* 

635 or NRC4 were co-expressed with *Rpi-amr1* or *Pp-Avramr1* on different knockout lines. *Rpi-*

636 *amr1-2273* or *Avramr1* alone were used as negative controls.

**Fig. S5:** Sequence exchange between *Rpi-amr1* homologs. Sequence exchange events were visually checked and highlighted (b and d) or identified by HybridCheck (a and c). For HybridCheck, sequence similarity was visualised using the colours of an RGB colour triangle (top); deviation from the default red, green and blue at positions with the same colour indicates regions where two sequences share the same polymorphisms, which is indicative of intra- or

642 inter-locus recombination. Line plot shows the percentage of SNPs shared at informative sites

643 between sequences in each of the three pairwise combinations for the triplet.

644 Fig. S6: Cloned *Rpi-amr1* CDS were mapped back to BAC\_5G using BLAT and visualized

on the BAC sequence using the Sushi package.

646

647 **Table S1:** Linked RAD markers identified based on tomato reference genome.

- **Table S2:** Molecular markers used in this study.
- **Table S3:** Amino acid sequence similarity between Rpi-amr1 homologs.
- **Table S4:** *S. nigrum* accessions used in this study.
- **Table S5**: Tajima's D analysis of *Rpi-amr1* and *Avramr1* homologs.
- **Table S6**: Evidence of sequence exchange between *Rpi-amr1* orthologs and paralogs from
- 653 SP2273 using 3SEQ.

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# 681 Supplementary Materials and Methods:

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#### 683 **Development of mapping populations**

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14 *P. infestans* resistant diploid *Solanum americanum* and relatives were used in this study (Table 1). The F1 populations were generated by crossing with a susceptible *Solanum americanum* accession 954750186 (working name SP2271) as a female parent. Heterozygous F1 progeny was allowed to self-pollinate to generate F2 segregating populations, or further back-crossed to the susceptible parent and allowed to self-pollinate until resistance to *P. infestans* co-segregated as a monogenic trait.

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# 692 *P. infestans* infection assay

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694 *P. infestans* isolates were cultured on rye and sucrose agar (RSA) medium at 18 °C for 10 days. 695 Sporangia were washed off with cold water and incubated at 4°C for 1-2 h to induce zoospore 696 release. Detached leaves were inoculated on the abaxial side with 10  $\mu$ l droplets of zoospore 697 suspension (50-100,000 per ml). The inoculated leaves were incubated at 18°C in high 698 humidity under 16 h day/8 h night photoperiod conditions. Disease scoring was done at 5-7 699 days after infection.

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# 701 **DNA and RNA extraction**

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RenSeq experiments (both short- and long-reads protocols) were conducted on gDNA freshly
extracted from young leaves using the DNeasy Plant Mini Kit (Qiagen) according to the
manufacturer's protocol. For the cDNA RenSeq experiment, RNA was extracted using TRIReagent (Sigma-Aldrich, MO, USA) and Direct-zol RNA MiniPrep Kit (Zymo Research, CA,
USA), following manufacturer's recommendations.

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# 709 Mapping of Rpi-amr1

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To map the underlying resistance gene from the resistant parent 954750184 (working name

- SP2273), we generated an F2 segregating population which was phenotyped with *P. infestans*
- isolates EC1\_3626 and 06\_3928A. Selected resistant plants were self-pollinated and up to 100

plants from F3 populations were screened for resistance and susceptibility with *P. infestans*isolates EC1\_3626 and 06\_3928A. gDNA from susceptible F2 and F3 plants (BS pool), as well
as gDNA from the resistant (R) and susceptible parent (S) were subjected to RenSeq using
Solanaceae bait library<sup>1</sup> and sequenced with Illumina GAII 76 bp paired-end reads. Preprocessing, assembly, mapping and SNP calling was performed as described earlier<sup>1,2</sup>.

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The same gDNA samples were used in a RAD-seq experiment using PstI digestion and Illumina HiSeq sequencing, which was outsourced to Floragenex Inc. (OR, USA). Bioinformatic analysis was also performed by Floragenex using *Solanum lycopersicum* genome as a reference<sup>3</sup>. SNP calling resulted in sixteen polymorphic sites with eleven of them locating at the top of chromosome 11 (Supplementary table 1). The remaining ones were randomly distributed on chromosomes 4 and 1.

726

We additionally outsourced Whole Genome Shotgun sequencing (WGS) of R and S samples to BGI (BGI, Shenzhen, China) for ~30 deep Illumina HiSeq sequencing with 100PE. Reads from the resistant parent were assembled as described in<sup>2</sup> and we used our previously published *in silico* trait mapping pipelines to perform SNP calling and detection of polymorphisms linked to disease resistance<sup>1,2</sup>. Contigs polymorphic between R and S parents were further aligned to

- the DM reference genome to identify their position.
- 733

734 Screening a set of markers derived from these three approaches on gDNA of 94 susceptible F2 735 and F3 plants identified 12 markers linked to resistance response that flank the *R* locus between 7.5 cM to one side and 4.3 cM to the other side (WGS, Table S1). Four of these markers were 736 737 found to co-segregate with the resistance, and two others located around 1 cM on either side, 738 CAPS marker RAD 3 to the distal side and the PCR marker WGS 1 to the proximal side 739 (Figure 1). Both 1 cM markers were subsequently used to genotype 1,793 F2 plants, and we 740 identified 228 recombinants (118 homozygous susceptible to one side and heterozygous to the 741 other, 110 homozygous resistant to one side and heterozygous to the other).

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The 118 informative recombinants (homozygous susceptible / heterozygous) were further genotyped using eight linked markers (Figure 1b), and tested in detached leaf assays for their response to *P. infestans* isolates EC1\_3626 and 06\_3928A. This revealed that markers CLC\_3 (WGS\_3) and RAD1 are flanking with a single recombination event for each marker, and CLC 2 (WGS 2), 56766 and 46418 are co-segregating with the resistance locus (Figure 1b).

748 Comparison of the linkage map (Figure 1) with the potato reference genome<sup>4</sup> identified the

homogeneous CNL-3 NLR gene sub-family to be within the co-segregating locus. This cluster
 comprises 18 members on potato reference chromosome 11.

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#### 752 BAC clones identification and analysis

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754 Construction and screening of 5x BAC library from resistant parent SP2273 was outsourced to 755 BioS&T company (Quebec, Canada). Two candidate BAC clones (5G and 12H) were 756 identified in PCR screen with WGS 2 marker-specific primers. BAC sequencing with RSII 757 PacBio platform and bioinformatic analysis was outsourced to Earlham Institute (Norwich, 758 UK); both BACs were assembled into single contigs with length of 125,327 bp (5G) and 759 144,006 bp (12H). While the co-segregating marker WGS 2 was present on both BAC clones, 760 a further co-segregating marker WGS 3 was only present on 12H. The BACs were further 761 assembled into one 212,773 contig (available in ENA under study number PRJEB38240). NLRs on the contig sequence were annotated using NLR-annotator<sup>5</sup> and Geneious 8.1.2 build-762 763 in ORF prediction tool. Gene models were annotated manually using cDNA RenSeq data 764 generated from S. americanum accession SP2273 as described below.

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#### 766 **3' RACE**

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Total RNA was extracted using RNeasy Plant Mini Kit (Qiagen) and treated with RNase-Free 768 769 DNase (Qiagen) following manufacturer's instructions. First strand cDNA was synthesized from total RNA using SuperScript<sup>™</sup> First-Strand Synthesis System for RT-PCR (Invitrogen, 770 771 CA, USA) with P7-oligoDT primer. The resulting product was amplified with P7- and gene 772 specific primers by using KAPA HiFi HotStart ReadyMix PCR Kit (Kapa Biosystems, Cape Town, SA) and cloned into pCR<sup>TM</sup>-Blunt II-TOPO vector by using Zero Blunt<sup>®</sup> TOPO<sup>®</sup> PCR 773 774 Cloning Kit (Invitrogen) and transformation was performed using One Shot<sup>™</sup> TOP10 775 Chemically Competent E. coli (Invitrogen). Isolation of plasmid DNA was performed with 776 NucleoSpin<sup>®</sup> Plasmid kit (MACHEREY-NAGEL, Duren, Germany).

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778 Resistance gene enrichment sequencing (RenSeq) and Gene enrichment sequencing779 (GenSeq)

SMRT RenSeq, short-read RenSeq and cDNA RenSeq were performed as described
 previously<sup>2</sup> and enriched libraries were sequenced at Earlham Institute, Norwich, UK (PacBio
 RSII, Illumina MiSeq) and Novogene, Hong Kong (Illumina HiSeq).

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784 Illumina GenSeq was performed as described above (Illumina RenSeq), except GenSeq baits<sup>6</sup>
785 were used instead of RenSeq baits.

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PacBio reads were processed and assembled using Geneious R8.1.8<sup>7</sup> as described<sup>2</sup>. NLR
 coding sequences were predicted with Geneious and AUGUSTUS<sup>8</sup> and annotated with NLR parser<sup>5</sup>.

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791 To infer linked polymorphisms, the quality control for Illumina paired-end reads was 792 performed using Trimmomatic<sup>9</sup> with standard settings. For the RenSeq, the paired reads were 793 mapped to PacBio-assembled contigs from the resistant parent, while GenSeq reads were 794 mapped to the reference DM genome (PGSC DM v4.03 pseudomolecules.fasta), using BWA mapper<sup>10</sup> with default settings. PCR duplicates and unmapped reads were removed and 795 Mpileup files to find out potential linked SNPs were created using SAMtools<sup>11</sup>. Mpileup files 796 were processed with VarScan<sup>12</sup> set to minimum read depth 20, minimum variant allele 797 frequency threshold 0.1, and minimum frequency to call homozygote 0.98. The candidate SNPs 798 were manually inspected using Savant genome browser<sup>13</sup>. TopHat<sup>14</sup> with default settings was 799 800 used to map cDNA Illumina reads to assembled PacBio data. All the tools used in this study 801 were embedded in The Sainsbury Laboratory (TSL) customized Galaxy instance, if not stated 802 otherwise.

#### 803 Transient complementation of a candidate genes in *N. benthamiana*

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The candidate genes were PCR amplified from gDNA with their own promoters (1-2 kb upstream of start codon) and up to 1 kb terminator elements, and cloned into USER vector as described<sup>2</sup>. Transient complementation assays followed by *P. infestans* inoculation were performed as described<sup>2</sup>.

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### 810 Stable transformation of susceptible potato cultivar Maris Piper

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812 Stable transgenic plants with constructs carrying Rpi-amr1-2272, Rpi-amr1-2273 or 813 *Rpi-amr1a* under the control of their native regulatory elements were created in the 814 background of potato cultivar Maris Piper as described previously<sup>15</sup>. At least 10 815 independent transgenic lines were generated for each construct and tested for the 816 presence of the transgene using gene specific primers. All positive *Rpi-amr1-2272* and 817 Rpi-amr1-2273 lines showed resistance in DLA with P. infestans isolate 88069, while 818 *Rpi-amr1a* transgenic plants were fully susceptible. Selected lines of *Rpi-amr1-2272* 819 and Rpi-amr1-2273 were tested in DLA with 19 additional P. infestans isolates (Table 820 2). WT Maris Piper plants were used as a negative control.

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#### 822 Generation of *NRC2/3* knockout *N. benthamiana*

NRC4 and NRC2/NRC3/NRC4 knockout N. benthamiana lines were described 824 previously<sup>16,17</sup>. Knocking out of NRC2/NRC3 in N. benthamiana were performed 825 826 according to the methods described previously<sup>16</sup>. Forward primers CHW sgNbNRCs and reverse primer JC sgrna R<sup>16</sup> were used to clone sgRNA2.1-4, sgRNA3.1-4 into 827 Golden Gate level 1 vectors for different positions. Constructs of sgRNAs targeting N. 828 829 benthamiana NRC2 and NRC3 were assembled into level 2 vector pICSL4723 together 830 (pICH47732::NOSp::BAR, with pICSL11017 Addgene no. 51145) and pICH47742::35S::Cas9<sup>18</sup>. Leaf discs of *N. benthamiana* were transformed with the 831 832 binary vector pICSL4723 containing the BAR selection marker gene, Cas9 expression 833 cassette, and sgRNAs targeting NRC2 and NRC3. T0 transgenic plants were selected in 834 the medium with phosphinothricin (2 mg/L) and then transferred into the soil. The 835 progeny of the transformants were genotyped using amplicon sequencing as described previously<sup>16</sup> (Fig. S6a). T3 populations from the selected T2 plants were used for 836

837 further experiments. NRC2/3 knockout line (nrc23 1.3.1) did not exhibit any growth 838 defects when compared to the wild type plants (Fig. S6b). 839 840 **Phylogenetic tree construction** 841 842 Phylogenetic tree was generated from protein sequences of the cloned Solanaceae R 843 genes obtained from NCBI. Full-length sequences were aligned using ClustalW 1.74<sup>19</sup> 844 and the alignments were imported to the MEGA7<sup>20</sup> to build a maximum-likelihood phylogenetic tree with Jones-Taylor-Thornton (JTT) substitution model and 100 845 846 bootstraps. 847 848 Evolutionary analyses of *Rpi-amr1* and *Avramr1* homologs 849 CDS were aligned using MUSCLE<sup>21</sup> as implemented in seaview<sup>22</sup> with and without 850 outgroup (the closest homologs from S. lycopersicum and P. capsici for Rpi-amrl and 851 852 Avramr1, respectively). Calculations of diversity statistics and the MacDonald-Kreitmann Test were executed through DNAsp5.0<sup>23</sup>; DAMBE<sup>24</sup> was used to rule out 853 854 saturation. For *Rpi-amr1* homologs, the calculations were preformed separately on 855 annotated full-length sequences as well as the individual domains. 856 We used 3SEQ<sup>25</sup> to identify break points in the aligned CDS. To confirm gene 857 858 conversion events in *Rpi-amr1*, we mapped the CDS back to the BAC 5G sequence 859 using BLAT (minScore 1500, minMatch 93)<sup>26</sup>. The resulting .psl files were converted 860 into .bed files using a custom R script, prior to visualization using the R package Sushi<sup>27</sup>. 861 862 HybridCheck 863 864 For each accession, FASTA files of all *Rpi-amr1e* orthologs or *Rpi-amr1* paralogs in combinations of three (triplets) were generated and aligned using MUSCLE v3.8.31<sup>21</sup>. 865

The sequence triplets were analysed using HybridCheck<sup>28</sup> to detect and date recombination blocks between *Rpi-amr1* orthologs (sliding windows = 200bp) or paralogs (sliding windows = 100 bp); non-informative sites were removed from the sequence triplets. Figures showing sequence similarity were plotted (MosaicScale = 50)

- 870 with HybridCheck and formatted using R v3.2.0 (https://www.r-project.org). The
- 871 colour of each sequence window was calculated based on the proportion of SNPs shared
- 872 between pairwise sequences at informative sites.

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