Identification of Natural Resistance Mediated by Recognition of *Phytophthora infestans* Effector Gene *Avr3aEM* in Potato

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Late blight is considered the most renowned devastating potato disease worldwide. Resistance gene (R)-based resistance to late blight is the most effective method to inhibit infection by the causal agent *Phytophthora infestans*. However, the limited availability of resistant potato varieties and the rapid loss of R resistance, caused by *P. infestans* virulence variability, make disease control rely on fungicide application. We employed an *Agrobacterium tumefaciens*-mediated transient gene expression assay and effector biology approach to understand late blight resistance of Chinese varieties that showed years of promising field performance. We are particularly interested in *PiAvr3aEM*, the most common virulent allele of *PiAvr3aKI* that triggers a R3a-mediated hypersensitive response (HR) and late blight resistance. Through our significantly improved *A. tumefaciens*-mediated transient gene expression assay in potato using cultured seedlings, we characterized two dominant potato varieties, Qingshu9 and Longshu7, in China by transient expression of *P. infestans* effector genes. Transient expression of 10 known avirulence genes showed that *PiAvr4* and *PiAvr8 (PiAvrsmira2)* could induce HR in Qingshu9, and *PiAvrvt1.1* in Longshu7, respectively. Our study also indicated that *PiAvr3aEM* is recognized by these two potato varieties, and is likely involved in their significant field performance of late blight resistance. The identification of natural resistance mediated by *PiAvr3aEM* recognition in Qingshu9 and Longshu7 will facilitate breeding for improved potato resistance against *P. infestans*.

**Keywords:** potato late blight, *Phytophthora infestans*, *PiAvr3aEM*, Qingshu9, Longshu7, *Agrobacterium tumefaciens*, RXLR effectors, hypersensitive response
INTRODUCTION

Potato (Solanum tuberosum L.) is regarded as the fourth-largest food crop and the main non-cereal crop worldwide which is influenced by the destructive and notorious late blight disease (Aguilera-Galvez et al., 2018). Phytophthora infestans is the causative agent that can destroy all potato parts, including leaves, stems and tubers (Fry, 2008), and is the main threat to potato production and responsible for 16% of yield losses globally (Haverkort et al., 2016). Similar to other crops, disease management using resistant varieties is one of the most effective strategies, environmentally and economically, to control late blight disease (Fry, 2016). Plant immunity is activated by detecting conserved microbial molecules, microbe (pathogen)-associated molecular patterns (MAMPs or PAMPs), known as pattern-triggered immunity (PTI), as well as by detecting the pathogen effectors, known as effector-triggered immunity (ETI) (Jones and Dangl, 2006). Plant pathogens can successfully colonize plant hosts by delivering effector proteins that repress host immunity and increase disease severity (Turnbull et al., 2017). In turn, few effectors might be recognized by the correspondent resistance (R) proteins, triggering a rapid immune response known as effector-triggered immunity (ETI), which often leads to an hypersensitive response (HR) cell death (Jones and Dangl, 2006; Turnbull et al., 2017).

To achieve effective control of late blight, potato breeders have to adopt novel techniques and strategies in various aspects such as detection and identification of new Rpi genes, their introgression and field application (Vleeshouwers and Oliver, 2014). Great efforts were made at the beginning of the last century to introgress Rpi genes into potato varieties from the wild Mexican species Solanum demissum to provide resistance to P. infestans in the cultivated potato S. tuberosum. This also led to the development of differential potato lines with 11 distinct recognition specificities, called R1–R11 (Black et al., 1953; Malcolmson and Black, 1966). Many R genes (Rpi) have been identified, cloned and some of them were introgressed into potato cultivars from wild Mexican Solanum species (Goodwin et al., 1992; Grunwald and Flier, 2005; Goss et al., 2014), including R1–R11, R3a, R3b, R9a and Rpi-demf1 from S. demissum (Ballvora et al., 2002; Huang et al., 2005; Lokossou et al., 2009; Jo et al., 2015), Rpi-blb1(RB), Rpi-blb2, Rpi-blb3, Rpi-abpt and Rpi-bt1 from S. bulbocastanum (Song et al., 2003; Van der Vossen et al., 2003; Park et al., 2005; Van der Vossen et al., 2005; Lokossou et al., 2009), Rpi-sto1,2, Rpi-pta1,2, and Rpi-plt1 from S. stoloniferum (Vleeshouwers et al., 2008; Wang et al., 2008; Champouret et al., 2009), Rpi-amr3 from S. amercianum (Witek et al., 2016), Rpi-mch1 from S. micoaocanum (Sliwka et al., 2012a), and Rpi1 from S. pinnatisectum (Kuhl et al., 2001; Lokossou et al., 2010). Additionally, further R genes have been identified in another center of genetic diversity of tuber-bearing Solanum, the Andean region in South America, such as Rpi-vnt1 from S. venturi (Foster et al., 2009; Pel et al., 2009), Rpi-mcq1 from S. mochiquense (Smilde et al., 2005), Rpi-ber from S. berthaultii (Rauscher et al., 2006), Rpi-mcd1 from S. microdontum (Tan et al., 2008), Rpi-pcs from S. paucissectum (Villamon et al., 2005), Rpi-capI from S. capsicrabactatum (Jacobs et al., 2010), Rpi-rzc1 from S. ruiz-cabellosi (syn. S. sparsipilum) (Śliwka et al., 2012b; Brylińska et al., 2015) and Rpi-chc1 from S. chacoense (Zhu et al., 2015).

Much attention has been paid to RXLR effectors since the cloned avirulence protein (Avr) of oomycete pathogens belong to this type of effector, such as P. sojae Avr1b (Shan et al., 2004), Hyaloperonospora arabidopsidis ATR13 (Allen et al., 2004) and ATR1 (Rehmany et al., 2005), and P. infestans PiAvr3a (Armstrong et al., 2005). So far, over 10 Avr genes, identified by 10 cognate-recognition Rpi genes, have been described in P. infestans, including PiAvr1 (Van der Lee et al., 2001; Ballvora et al., 2002), PiAvr2 (Gilroy et al., 2011), PiAvr3a (Armstrong et al., 2005), PiAvr3b (Rietman et al., 2012), PiAvr4 (Van Poppel et al., 2008), PiAvr8 (Vossen et al., 2016), PiAvr1bl1 (Song et al., 2003; Vleeshouwers et al., 2008), PiAvr2bl2 (Van der Vossen et al., 2005; Oh et al., 2009), PiAvrvt1 (Foster et al., 2009; Pel et al., 2009), PiAvrSmira1 and PiAvrSmira2 (Rietman et al., 2012; Yoshida et al., 2013) as shown in (Supplementary Table S1). P. infestans is predicted to encode 563 RXLR effector genes which were mainly found located in repeat-rich or gene-sparse regions of the genome, meaning that more rapid evolution compared to other genes located in gene-dense regions (Haas et al., 2009; Yin et al., 2017). It might explain its ability to escape host defense mechanisms (Lenman et al., 2016). It has been well-documented that RXLR effectors play important roles in potato–P. infestans interactions. In rare cases of recognition by the cognate R genes, they mediate late blight resistance by triggering HR; in most cases, they are not recognized and function as typical virulence factors by interfering with host cell structure and function, resulting in enhancing plant susceptibility (Huang et al., 2019).

There are generally two strategies to improve late blight resistance. The first is the deployment of many different R genes to offer tentative durable resistance since changes of multiple effectors are predicted to increase the penalty. The second strategy is to identify genes that are capable of recognizing various effectors or core effectors. In fact, the identified R genes from varieties that showed durable disease resistance were confirmed to be able to recognize two or more effectors. For example, the potato Rpi-blb1, known as RB (Song et al., 2003) recognizes PiAvrblb1, ipiO1 and ipiO2 (Vleeshouwers et al., 2008; Chen et al., 2012); the Rps1k of soybean recognizes two P. sojae effectors, Avr1b and Avr1k (Shan et al., 2004; Dou et al., 2010).

Yet, P. infestans PiAvr3a is a well-characterized P. infestans RXLR effector that is highly expressed during the biotrophic phase of infection [2–3 days post in filtration (dpi)] (Haas et al., 2009; Chaparro-Garcia et al., 2015). PiAvr3a is essential for full virulence, pathogenicity and suppression of host immunity, including PTI and ETI, by suppressing the programmed cell death (PCD) triggered by the elicitor NIN1, a secreted P. infestans protein with PAMP properties, by interacting with and stabilizing the host U-box E3 ligase CMPG1 (Bos et al., 2006; González-Lamothe et al., 2006; Bos et al., 2010; Gilroy et al., 2011), as well as targeting the receptor-mediated endocytosis dynamin-related protein 2B (DRP2B), clathrin-mediated...
endocytosis (CME) (Chaparro-Garcia et al., 2015). Two major allelic isoforms of PiAvr3a have been identified in *P. infestans* populations that have a difference in three amino acids in mature protein positions 19, 80 and 103 (Chapman et al., 2014). Avr3a (S19) K80M103 is known as PiAvr3aEM while Avr3a (C19) K80I103 is known as PiAvr3aKI (Vleeshouwers et al., 2011; Yang et al., 2018). Unlike PiAvr3aEM, PiAvr3aKI activates the potato R3a resistance protein to trigger ETI and confers avirulence to heterozygous or homozygous strains of the pathogen (Armstrong et al., 2005; Chaparro-Garcia et al., 2015). Therefore, *P. infestans* isolates, expressing only the PiAvr3aEM variant, can evade R3a recognition and do not trigger HR (Armstrong et al., 2005; Bos et al., 2006). Identification of natural R genes that can recognize PiAvr3aEM are promising approaches to improve late blight resistance (Bos et al., 2010; Segretin et al., 2014).

*Agrobacterium tumefaciens*-mediated transient gene expression technology is a rapid, widely and easily performed assay that is commonly used in gene expression analysis and functional genomics studies in many plant species, including *Arabidopsis thaliana*, tobacco, tomato, soybean, citrus, grapevine and potato (Vleeshouwers et al., 2008). Typically, *A. tumefaciens*-mediated transient expression assays can be utilized for several purposes, such as i) functional genomics tools for transient overexpression of a gene in planta, ii) reverse genetic studies of a gene by virus-induced gene silencing (VIGS) or RNA interference (RNAi) technology, iii) rapid accessible production of recombinant proteins, iv) pathogen effector assays for the genetic components of the selected cultivars disease resistance.

In this study, we utilized the optimum conditions for *A. tumefaciens*-mediated transient assays in potato and performed analyses of two potato varieties for their capability to recognize a set of *P. infestans* known effectors, as part of our effort in understanding late blight resistance of potato varieties that showed promising field performance. This led to the identification of natural resistance, mediated by recognition of *P. infestans* Avr3aEM, which will facilitate potato breeding for improved late blight resistance.

**MATERIALS AND METHODS**

**Plant Materials and Growth Condition**

Qingshu9 and Longshu7 are dominant potato varieties in Northwestern China. Qingshu9 was derived from crosses of two parents “387521.3 × APHRODITE”, while Longshu7 was derived from Fedori×Zhuangshu3. Potato cuttings have been cultured in a sterilized MS medium for four weeks (Murashige and Skoog, 1962). Next, the germinated seedlings were transferred for another four weeks into vermiculite, and then planted in pots that contain a mix of sterilized vermiculite and peat moss (V/V = 1:2). Also, some potato differentials, including R1, R2, R3a, R4 and R8, were used for the optimization of agroinfiltration assays. In addition, nine potato breeding lines were studied and agro-infiltrated with *Pi*Avr3aEM. Progeny lines resulting from crossing Qingshu9 with Qingshu2, ND, NSS1-5, and Jizhang8, respectively, and Longshu7 with CIP01, CIP03, CIP16, CIP30, and CIPLO6408, respectively, were evaluated. At least 20 progenies from each cross were evaluated. Potato plants were grown under standardized conditions in a greenhouse within a temperature range of 18–22°C and under a day/night regime of 16 h/8 h. Fully-expanded leaves of the 4-week-old seedlings were used for infiltration with bacterial cell suspensions of *A. tumefaciens* strain AG1L1 that carry a number of *Avr* genes to be evaluated.

**Cloning and Vector Construction of *P. infestans* Avr Genes**

All tested *Avr* genes were amplified from their plasmid DNA previously constructed into pK7WG2 vector, using TransStart® FastPfu DNA Polymerase (Applied Biosystems, USA) with *Avr* genes-specific primers containing the restriction enzyme recognition sites as shown in Supplementary Table S2. The PCR amplicons were purified using the TIANGEN Universal DNA Purification Kit (TianGen Biotech Co., Ltd., Beijing, China). The purified amplicons and the pART27 cloning vector were digested with the corresponding restriction enzymes and ligated together using T4 DNA ligase (Promega, USA). The ligation mixtures were transferred to *E. coli* DH5α competent cells by electroporation using standard protocols. Transformed colonies were cultured on LB medium supplemented with 100 µg ml⁻¹ of spectinomycin and incubated at 37°C. Positive clones were confirmed by sequencing. The confirmed plasmid constructs were then transformed into *A. tumefaciens* strain AG1L1 by the heat shock method. The transformed cell cultures were applied to LB plates containing antibiotics (100 µg ml⁻¹ of spectinomycin, 20 µg ml⁻¹ of rifampicin) and placed in a 28°C incubator for 2 days. A single colony was transferred using sterilized toothpicks to the liquid LB broth having the same antibiotics and incubated at 200 rpm in a shaker at 28°C for 2 days.

**Transient Agro-Infiltration Assays**

The optimized conditions of agroinfiltration-mediated transient expression assays were 3–4 or 9–10 week-old potato seedlings, *A. tumefaciens* strain AG1L1 and OD₆₀₀ value of 0.4. *A. tumefaciens* cells were grown in LB medium (supplemented with 50 µg ml⁻¹ of gentamicin, 20 µg m⁻¹ of rifampicin and 100 µg m⁻¹ of spectinomycin, 20 µg ml⁻¹ of rifampicin, respectively) up to the log phase of development. The bacterial solution was then centrifuged at room temperature (20°C, 4,000g, 3 min), followed by resuspension in an inducing media (10 mM MES, 200 µM acetosyringone, 10 mM MgCl₂, pH 5.6). The optical density of the *A. tumefaciens* suspensions was adjusted to OD₆₀₀ value of 0.4 and incubated before infiltration for 1–3 h at room temperature. Agroinfiltration experiments were carried out at room temperature 20 ± 2°C (Dillen et al., 1997; Su et al., 2012) on potato seedling leaves 4-week-old and the results were scored from 3 dpi and typically photographed at 5–7 dpi.
Optimization of Agro-Infiltration Assays

To evaluate factors affecting the agro-infiltration assay, three parameters were assessed, including different *A. tumefaciens* strains (AGL1 and GV3101), bacterial cell densities (OD\textsubscript{600} values of 0.2, 0.3, 0.4, 0.5, and 0.6), and different growth ages of cultured potato plants (3–4, 6–7 and 9–10 week-old). All experiments have been repeated three times with 20 replicates for each. In our study, potato seedling age refers to the time starting from tissue culture seedlings transferred to the soil matrix, after acclimatization in the vermiculite, to the time of experimentation.

**Effectors Screen**

Ten *P. infestans Avr* effectors were investigated including, *PiAvr1, PiAvr2, PiAvr3a\textsuperscript{KI}, PiAvr3b, PiAvr4, PiAvrb1b, PiAvrb2b, PiAvrsmira1, PiAvr8 (PiAvrsmira2) and PiAvrvt1.1* (Supplementary Table S1). *A. tumefaciens* strain AGL1 carrying each of these effectors was used for infiltration in the two varieties with a concentration of OD\textsubscript{600} value of 0.4. Further investigation was done for *PiAvr3a* alleles, *PiAvr3a\textsuperscript{EM} and PiAvr3a\textsuperscript{KI}*. Forty leaves were agro-infiltrated for each *Avr* effector in eight independent experiments with five replicates for each. All pictures were taken 5–7 days later of infiltration.

**Detection of R8 and Rpi-vnt1 by PCR Amplification**

Genomic DNA was isolated from fresh leaf tissue of both potato varieties using the CTAB-based protocol. The genomic DNA was then subjected to PCR with primers specific to R8 and Rpi-vnt1 (Foster et al., 2009; Pel et al., 2009; Vossen et al., 2016), as listed in the supplementary Table S3. PCR reactions were performed using FastPfu DNA polymerase (Applied Biosystems, USA). Each PCR reaction contained 30 μl PCR mix, including 6 μl 5X FastPfu Buffer, 2.4 μl dNTPs (0.2 mM), 1 μl total genomic DNA (100 ng), 0.2 μl MgSO\textsubscript{4} (50 mM), 0.6 μl each forward and reverse primers (0.2 mM), 0.6 μl FastPfu DNA polymerase (2.5 units) and 18.6 μl dH\textsubscript{2}O. The PCR amplification was carried out by denaturing at 95°C for 2 min, followed by 40 cycles of 94°C for 20 s, 55°C for 20 s and 72°C for 1 min, and a final extension time of 5 min at 72°C. PCR products were separated by gel electrophoresis on a 1% agarose gel and DNA bands were visualized under UV on the Quantum CX5 Imaging System.

**P. infestans Infection Assays on Detached Potato Leaves**

*P. infestans* isolates were cultured and maintained on a rye sucrose agar (RSA) medium. All plates were then grown at 16°C in darkness for two weeks. The sporangial suspensions were prepared by washing and rubbing the culture with 5 ml distilled water. Then, the sporangial concentration was adjusted to 4 × 10\textsuperscript{5} sporangia/ml before cooled down for 2 h at 4°C to promote release of motile zoospores for inoculation (Tian et al., 2015). Leaflets of 6–10 week-old potato plants were placed abaxially on plastic trays on a filter paper saturated with dH\textsubscript{2}O. All leaflets were drop-inoculated with 15 μl sporangial/zoospore suspension on the abaxial side. Six *P. infestans* isolates, PJY009, PJY048, PJY061, Pa21106, Pd21410 and F48, were used in the inoculation assays (Supplementary Figure S2). Inoculation with dH\textsubscript{2}O was considered as a control treatment. All plastic trays were covered by a plastic wrap and incubated in a growth chamber at 16–18°C with >75% relative humidity in the darkness in order to ensure infection. Results were recorded as a lesion diameter of the inoculated area were and pictures were taken five days after inoculation. Disease resistance or susceptibility were recorded by using a scale reported for disease severity (Sun, 2012).

**RESULTS**

**Optimization of *A. tumefaciens*-Mediated Transient Gene Expression Assay**

The outcome of plant-*Agrobacterium* interactions is determined by the genetic background of both partners. In addition to the efficiency of transient gene expression, the frequent non-specific necrotic response is a major concern in the use of this assay. We therefore examined for suitable *A. tumefaciens* strains with reduced background necrotic reaction in potato. Six different *A. tumefaciens* strains, *Agro*-1D124g, GV3101, AGL1, 1100, LBA4404, and EHA105, were evaluated on ten different potato varieties (Data not shown). Even though the OD\textsubscript{600} value was very low, strains 1100, LBA4404 and EHA105 induced a high rate of background necrosis on most of the potato cultivars. However, with a lower concentration of bacterial suspensions, GV3101 and AGL1 strains showed a significant reduction of background reaction on most potato varieties. Thus, strains GV3101 and AGL1 were employed to further investigate their transient expression efficiencies on the potato. The efficiency assay was examined by the HR symptoms that resulted from the co-infiltration of *P. infestans Avr* gene *PiAvrb1b* and its cognate resistance gene *R8*. The results showed that the AGL1 strain was more efficient than GV3101 in terms of triggering specific HR mediated by co-expression of *RB* and *PiAvrb1b* (Figure 1).

We also evaluated various potato seedling growth ages (3–4, 6–7 and 9–10 week-old) for the effect on the efficiency of the agro-infiltration assay. Five potato differential lines, including *R1, R2, R3a, R4* and *R8*, were examined and agro-infiltrated with *A. tumefaciens* AGL1 suspensions (OD\textsubscript{600} of 0.4) carrying the *P. infestans Avr* genes *PiAvr1, PiAvr2, PiAvr3a\textsuperscript{KI}, PiAvr4*, and *PiAvr8*, respectively. The positive control was the co-infiltration of mixed agrobacteria carrying *PiAvrb1b* and *RB* which would lead to HR while the negative control was the GFP. The most consistent and efficient infiltration was observed while using terminal leaflets from 3–4 and 9–10 week-old potato plants in all tested differential lines, while the 6–7 week potato leaves exhibited less efficient transient expression (Supplementary Figure S1). We speculated that the potato leaves were younger in 3–4 weeks when the leaves have just spread and the main veins were developed, but the lateral veins were not obvious, allowing easier infiltration in whole leaves. Meanwhile, the leaves of 9–10 week-old seedlings were fully developed, and the main and lateral veins were well developed, allowing efficient infiltration.
between the two lateral veins. However, the main veins and lateral veins of leaves of 6–7 week-old seedlings were all developed, still small interveinal spaces on the abaxial side hinder the infiltration process, making the bacterial solution restricted to a fixed grid, necessitating more infiltration sites. The optimum condition was utilized for further analysis which could be summarized as using the A. tumefaciens strain AGL1, with an OD\textsubscript{600} value of 0.4 and leaves of the 3–4 or 9–10 week-old seedlings.

To further confirm our improved agroinfiltration assay, we examined known Avr effector genes for their capability in triggering genotype-specific HR. A. tumefaciens AGL1 bacterial suspensions carrying the P. infestans Avr genes PiAvr1, PiAvr2, PiAvr3a\textsuperscript{KI}, PiAvr4, and PiAvr8 were infiltrated in potato differential lines carrying genotype-specific R genes R1, R2, R3a, R4, and R8, respectively. Each density (OD\textsubscript{600} values of 0.2, 0.4 and 0.6) showed a different level of transient expression. The bacterial suspensions with OD\textsubscript{600} value of 0.4 consistently displayed the highest efficiency in bacterial infiltration assays, as all tested Avr genes induced genotype-specific HR in all tested differential lines (Figure 2). While at a higher agrobacterial concentration of OD\textsubscript{600} of 0.6, an increase of HR response for all tested Avr genes and significant background necrosis for the negative control of GFP expression were observed in all tested differentials, though at the lower agrobacterial concentration (OD\textsubscript{600} of 0.2) the HR triggered by PiAvr3a and PiAvr8 in potato differential lines R3a and R8, respectively, were not visible.

**Evaluation of Two Potato Varieties for Recognizing Known P. infestans Avr Genes**

To understand late blight resistance of two Chinese potato varieties, Qingshu9 and Longshu7, that showed excellent field performance with a low percentage of disease incidence and severity (Wang et al., 2018), we evaluated whether they contain known R genes by examining their capability to recognize corresponding 10 P. infestans Avr genes. Both Qingshu9 and Longshu7 showed typical genotype-specific HR phenotypes 5 days post infiltration with A. tumefaciens AGL1 cell suspensions with an OD\textsubscript{600} of 0.4. Qingshu9 showed HR triggered by two P. infestans Avr genes, PiAvr4 and PiAvr8, suggesting the presence of R genes R4 and R8. Longshu7 showed HR triggered by
PiAvrvnt1.1, indicating the existence of Rpi-vnt1 (Figure 3). Furthermore, the presence of R8 and Rpi-vnt1 in Qingshu9 and Longshu7, respectively, was preliminarily analyzed by PCR amplification using gene-specific primers (Foster et al., 2009; Pel et al., 2009; Vossen et al., 2016) (Supplementary Figure S3). However, whether they are functional R genes needs further validation. PCR amplification might provide possibility for their presence since it is highly dependent on the specific primers, while the potential presence of functional R gene homologs/alleles may lead to false negative results. The agroinfiltration assay using effector genes is an efficient method to detect the presence of functional R genes, such as R8 and Rpi-vnt1.1 in this research.

Qingshu9 and Longshu7 Showed Genotype-Specific HR Triggered by Avr3aEM

P. infestans Avr gene PiAvr3aEM can be specifically recognized by the cognate R3a. However, the number of its virulent alleles, that escaped recognition by R3a, is very limited and the virulent allele PiAvr3aEM is widely present around the world, suggesting the vital role of PiAvr3a in P. infestans pathogenesis. The identification of varieties with capable PiAvr3aEM recognition that make it possible for breeding new varieties with the capability to recognize both PiAvr3aKI and PiAvr3aEM, which is predicted to improve durable resistance against late blight. Both Qingshu9 and Longshu7 showed an HR upon PiAvr3aEM infiltration, but not upon PiAvr3aKI (Figure 4).

Recognition of P. infestans Avr3aEM by Qingshu9 and Longshu7 Is Likely Conferred by a Single Gene

Given the fact that PiAvr3aKI (Armstrong et al., 2005) is recognized by R3a (Huang et al., 2005), a cloned and well-studied R gene, we predict that PiAvr3aEM is similarly recognized by a single R gene. We therefore employed several independent F1 segregation populations to ensure that the PiAvr3aEM recognition is conditioned by a single R gene, by examining whether the PiAvr3aEM recognition-triggered cell death segregates. We performed agroinfiltration assays for progenies derived from a
total of nine crosses for the two responsive varieties, with five crosses using Longshu7 as a resistant parental with five non-responsive potato clones as the susceptible parental, including CIP01, CIP03, CIP16, CIP30, and CIPL06408. Qingshu9 as the resistant parental was crossed with four non-responsive potato clones as the susceptible parental, including Qingshu2, ND, NSS1-5, and Jizhang8. Twenty F1 progenies from each cross were tested for their response upon infiltration with PiAvr3aEM, with a total of 30 infiltration sites for each progeny. The results showed that progeny derived from two investigated Longshu7 crosses showed an HR response upon PiAvr3aEM infiltration at a rate of 1:1 for each cross, including Longshu7 X CIP01 and Longshu7 X CIP16 as shown in (Figure 5) and supplementary (Supplementary Table S4). While for Qingshu9 crosses, progenies from only one investigated cross (Qingshu9 X ND) showed an HR response with a rate of 1:1 (Figure 5, Supplementary Table S4). Although we did not perform comprehensive genetic analysis, the segregation of PiAvr3aEM recognition strongly suggests that PiAvr3aEM recognition by Qingshu9 and Longshu7 is conditioned by a single R gene. Also, the results suggest that the R genes for PiAvr3aEM recognition in Qingshu9 and Longshu7 were heterozygous, and most, if not all, parental lines crossed with Qingshu9 or Longshu7 were unable to recognize PiAvr3aEM. Another possibility is that a helper/sensor NLR might be required for R3a function to initiate the immune signaling, resulting in an HR response, similar to the case of NRC4, a helper NLR essential for immunity triggered by Rpi-blb2 (Wu et al., 2017).

**DISCUSSION**

A critical step in the successful agroinfiltration-mediated transient expression is the establishment of harmonious interaction between the plant and *A. tumefaciens*. We also considered potential background non-specific necrotic reactions frequently caused by molecules from *A. tumefaciens*. We examined multiple *A. tumefaciens* strains (Agro-1D124g, GV3101, AGL1, 1100, LBA4404, and EHA105), potato genotypes, growth stages, and bacterial densities. This led to the conclusion that AGL1 was the most efficient strain with fewer background effects than GV3101, which is consistent with a report showing that strain AGL1 was preferred for potato whereas GV3101 was more suitable for *Nicotiana benthamiana* (Du et al., 2014).

Potato leaves at various growth stages were also a major concern and we found that the maximum level of bacterial infiltration was observed at the terminal leaflets from 3–4 to 9–10 week-old potato seedlings. The efficiency of infiltration became much lower when leaves were used from 6–7 week-old seedlings. Changes in the expression levels may be related to general changes in leaf physiology, especially soluble protein concentration during leaf aging (Halfhill et al., 2003; Wydro et al., 2006). Also leaf morphological characteristics including, leaf surface, the thickness of the cuticle layer and epidermis, stomatal size and frequency, veins, trichomes, the midrib structure, and interveinal distribution on the abaxial side leaves may affect infiltration efficiency (Abdullah and Halterman, 2018). Some developmental ages have a low density of trichomes on the abaxial side and the leaf veins on the surface are not prominent and as a result, facilitate infiltration (as 4-week-old leaves). Besides, the older leaves (9–10-week-old) where interveinal space has increased showed an increase in transient expression. Meanwhile, the 6–7-week-old leaves exhibited an irregular leaf surface with a high density of trichomes on the abaxial side, prominent veins and small interveinal spaces that can hinder the infiltration process.

Ten Avr genes had been described in *P. infestans* and their gain-of-virulence alleles were reported (Vleeshouwers and Oliver, 2014). In this study, we aimed at detecting and identifying R genes in potato varieties that showed excellent performance against late blight. Agroinfiltration analysis of 10 *P. infestans* Avr genes was done using *A. tumefaciens* strain AGL1 with OD600 value of 0.4 to

![FIGURE 5 | Segregation of HR induced by recognition of *P. infestans* PiAvr3aEM in F1 populations using Longshu7 or Qingshu9 as a parental. Progenies derived from crosses Longshu7 X CIP01 and Longshu7 X CIP16 showed typical HR as that in Longshu7, and progenies from cross Qingshu9 X ND showed typical HR as that in Qingshu9. GFP was used as negative control and co-expression of PiAvrblb1 and RB was used as a positive control. The number of HR-responsive progenies/total number of progenies, were indicated in each crosses, with each progenies examined with 30 infiltration sites. All pictures were taken at 5–7 dpi.](image)
evaluate two potato varieties, Qingshu9 and Longshu7, whether they encode R genes that may recognize these Avr genes. The agroinfiltration in Qingshu9 resulted in activating a typical HR towards three effector genes including, PiAvr4, PiAvr8 and PiAvr3aEM, which is a virulent allele of PiAvr3a that can be recognized by R3a, whereas Longshu7 exhibited a typical HR by two P. infestans effector genes, PiAvr3aEM and PiAvrSmira1. According to the gene-for-gene hypothesis, Avr genes are detected by its counterpart R genes (Anderson et al., 2015; Yin et al., 2017). These results suggest that Qingshu9 may carry at least three R genes, including R4, R8, and R3a*, whereas Longshu7 carries at least two R genes, Rpi-vnt1 and R3a*.

PiAvr3a was highlighted and has been extensively studied which is expected to be a useful target for potato breeders seeking durable resistance (Cooke et al., 2012). PiAvr3a appears to be a core effector of P. infestans since it’s among few effectors that are conserved across several Phytophthora species and it is consistently induced within the early stages of P. infestans infection (Yin et al., 2017). Besides, it is involved in the suppression of PTI and ETI (Gilroy et al., 2011; Franco-Orozco et al., 2017). So far, there are only two detected PiAvr3a alleles among P. infestans populations (Bos et al., 2010). The avirulent allele, PiAvr3aEM, is recognized by R3a, while its virulent allele, PiAvr3aEM, evades recognition by R3a (Armstrong et al., 2005; Chapman et al., 2014).

Previous studies showed successful recognition of PiAvr3aEM by engineering potato resistance gene R3a (Chapman et al., 2014) and by screening the R3a variants library resulting from random mutagenesis of the full-length R3a coding sequence (Segretin et al., 2014). Remarkably, our results offer a new natural resistance gene that can recognize PiAvr3aEM in two potato varieties, suggesting that these two varieties are potentially undergoing R3a*-mediated recognition responses. Both varieties were derived from crosses that used different parents, for Longshu7 being derived from FederalxZhuangshu3, while Qingshu9 from 387521.3 × APHRODITE, suggesting that both varieties might contain a functional homolog of the R3a* resistance gene and both are very likely heterozygous. It’s also possible that they might have two different forms of R3a* that mediate PiAvr3aEM recognition. Both varieties didn’t show any PiAvr3KI recognition, suggesting that they don’t carry the known R3a.

Further work on the survey of PiAvr3EM-meditated HR on progenies derived from crosses using either Qingshu9 or Longshu7 as a parental indicated that progenies from two Longshu7 crosses were detected with PiAvr3aEM-mediated HR, while progenies from a single Qingshu9 cross were detected for inducing an HR response, suggesting that the recognition of PiAvr3aEM is most likely conditioned by a single R gene R3a* in both varieties. Most lines that were crossed with either Longshu7 or Qingshu9, if not all, do not carry R3a*. The lack of PiAvr3aEM response in some populations is likely resulted from the heterologous nature of PiAvr3aEM recognition in the resistant parents and short of PiAvr3aEM recognition in the other parents. It’s also possible that we examined limited number of progenies. However, whether the PiAvr3aEM recognition in Longshu7 or Qingshu9 mediates late blight resistance needs additional pathogenicity tests. Under favorable infection conditions using detached leaves, our preliminary infection assays with diverse virulent P. infestans strains showed generally high levels of late blight resistance for Longshu7 and Qingshu9, though certain level of susceptibility was notable to several virulent strains (Supplementary Figure S2). There are potentially complicated interactions between effectors in suppression and triggering immune response. A promising efficient strategy to enhance late blight resistance is to integrate R3a that recognizes PiAvr3aKI and R3a* that mediates PiAvr3aEM response. However, whether such simple R gene combination is correlated with predicted enhanced durable late blight resistance needs confirmation by field assessments.

Rpi-Smira2 (R8) confers quantitative resistance under field conditions and associates with PiAvrSmira2 (PiAvr8) (Rietman et al., 2012; Hajianfar et al., 2014). In our study, Qingshu9 exhibited an HR response upon PiAvr8/PiAvrSmira2 infiltration, suggesting the presence of R8/Rpi-smira2 in Qingshu9. The PiAvr8/PiAvrSmira2-triggered HR in Qingshu9 was consistent with a previous report in which R8 is correlated with quantitative resistance and PiAvr8/PiAvrSmira2 triggered R8-mediated resistance (Rietman et al., 2012). Our findings are also consistent with a report in which genotype-specific HR was induced after R8-PiAvr8 co-infiltration as well as R8-like co-infiltration with PiAvr8 (Jiang et al., 2018). Notably, the NB-LRR gene R8 has been cloned and was thought to provide broad-spectrum and durable field resistance against P. infestans (Vossen et al., 2016; Jiang et al., 2018). It has been reported that Rpi-Smira2 co-localized with the R8 locus and both loci conferred similar resistance levels (Jo, 2013; Stefańczyk et al., 2017). Hence, it was suggested that Rpi-Smira2 and R8 are identical or functional homologs (Jo et al, 2011). In addition, many P. infestans isolates carry PiAvr8 that was reported to trigger an HR response of the R8 gene in disease resistant potato varieties and lines, such as Sarpo Mira from Europe, PB-06, S-60, and QTL dpIP09c from China, and Jacqueline Lee from USA (Vossen et al., 2016; Jiang et al., 2018), suggesting its vital role in the pathogen and the effectiveness of R8.

PiAvrvt1 is recognized by the potato resistance gene Rpi-phula/Rpi-vnt1 (Foster et al., 2009). Because of its polymorphism, it is associated with a response to a diversified target protein or recognition avoidance (Pel et al., 2009; Pais et al., 2017). Longshu7 showed HR toward Avrvnt1,1, suggesting that it carries the functional Rpi-vnt1 gene which may provide a high level and wide-spectrum late blight resistance (Stefańczyk et al., 2018).

In summary, we developed and used the optimized A. tumefaciens-mediated transient expression assays to evaluate two potato varieties Qingshu9 and Longshu7 that showed years of promising field late blight resistance for R genes they might carry, by detecting the presence of HR triggered by 10 known P. infestans Avr genes. This led to the identification of natural resistance mediated by recognition of PiAvr3aEM, a globally present virulence allele of PiAvr3aEM that plays vital roles in potato-P. infestans interactions. Interestingly, cloning and analysis of R3a* that mediates PiAvr3aEM recognition and other detected R genes in
Qingshu9 and Longshu7 will be interesting to make both good use of late blight resistance and improved understanding of disease resistance in future. Together with the identification of additional complementary R genes in the two varieties, these findings will facilitate the development of potato lines with a high level of late blight resistance, by pyramiding these promising R genes.

DATA AVAILABILITY STATEMENT

The datasets generated for this study are available on request to the corresponding author.

AUTHOR CONTRIBUTIONS

WS and YM designed the experiments. AE, JL, XW, CZ, and YM performed the experiments. AE, JL, YM, GW, JW, HL-K, and WS analyzed the data. AE, YM, and WS wrote the manuscript with contribution from all authors.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2020.00919/full#supplementary-material

SUPPLEMENTARY TABLE S1 | List of Avr genes of late blight pathogen P. infestans with their corresponding R genes.

SUPPLEMENTARY TABLE S2 | List of all specific primers used for cloning P. infestans Avr genes.

SUPPLEMENTARY TABLE S3 | List of specific primers used for detection of R8 and Rpi-vnt1 by PCR amplification.

SUPPLEMENTARY TABLE S4 | Segregation for PaAvr3aEM response in the F1 populations of crosses Longshu7 X CIP01, Longshu7 X CIP16, and Qingshu9 X ND.

SUPPLEMENTARY FIGURE S1 | Effect of various growth ages of potato seedlings on the efficiency of Agrobacterium tumefaciens-mediated transient expression. Leaves of three different growth stages, 3–4, 6–7 and 9–10 week-old seedlings, were investigated. Five P. infestans Avr genes (Avr1, Avr2, Avr3, Avr4, and Avr8) were expressed by A. tumefaciens AGL1, OD600 of 0.4, in five potato differential genotypes carrying cognate R genes R1, R2, R3a, R4 and R8, respectively. Co-infiltration of PaAvr8 in R and RB was used as a positive control, whereas GFP was used as a negative control. The number of HR sites/total number of infiltration sites were indicated in each treatment. All photographs were taken 5 dpi.

SUPPLEMENTARY FIGURE S2 | Infection of Qingshu9 and Longshu7 to different P. infestans strains. Detached leaves of 9–10 week-old plants were drop-inoculated with 4×10^4 sporangia/mL of six P. infestans strains with known virulence spectrum, including PY009, PY048, PY061 (Yunnan, Northwestern China), Pa21106, Pd21410 (Ningxia, Northwestern China) and F48 (Fujian, Southeastern China), with sterilized dH2O as the control treatment. Qingshu9 was highly resistant (R) to Pa21106, Pd21410 and F48, but susceptible (S) to PjY009, PY048 and PY061. Longshu7 was resistant to PY061, Pa21106, Pd21410 and F48, but susceptible to PY009 and PY048. Disease index (DI) were scored and photographs were taken 5 dpi. The experiment was repeated four times with five replications.

SUPPLEMENTARY FIGURE S3 | Detection of R8 and Rpi-vnt1 in Qingshu9 and Longshu7 by PCR amplification using gene-specific primers. Two sets of gene-specific primers for R8 and Rpi-vnt1 each were used for their detection.
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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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