Single Amino Acid Mutations in the Potato Immune Receptor R3a Expand Response to Phytophthora Effectors

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Both plants and animals rely on nucleotide-binding domain and leucine-rich repeat-containing (NB-LRR or NLR) proteins to respond to invading pathogens and activate immune responses. How plant NB-LRR proteins respond to pathogens is poorly understood. We undertook a gain-of-function random mutagenesis screen of the potato NB-LRR immune receptor R3a to study how this protein responds to the effector protein AVR3a from the oomycete pathogen Phytophthora infestans. R3a response can be extended to the stealthy AVR3aEM isof orm of the effector while retaining recognition of AVR3aK1. Each one of eight single amino acid mutations is sufficient to expand the R3a response to AVR3aEM and other AVR3a variants. These mutations occur across the R3a protein, from the N terminus to different regions of the LRR domain. Further characterization of these R3a mutants revealed that at least one of them was sensitized, exhibiting a stronger response than the wild-type R3a protein to AVR3aK1. Remarkably, the N336Y mutation, near the R3a nucleotide-binding pocket, conferred response to the effector protein PcAVR3a4 from the vegetable pathogen P. capsici. This work contributes to understanding how NB-LRR receptor specificity can be modulated. Together with knowledge of pathogen effector diversity, this strategy can be exploited to develop synthetic immune receptors.

Plant diseases represent a significant threat to human welfare, resulting in economic hardship, malnutrition, famine, and environmental degradation. Among the diverse plant pathogens, filamentous eukaryotic microorganisms such as oomycetes and fungi cause the most destructive crop diseases and are a recurrent threat to world food security (Fisher et al. 2012). The oomycete genus Phytophthora (“plant destroyer” in Greek) includes the most devastating pathogens of dicot plants (Fry 2008). More than 100 species of Phytophthora are known, including the notorious Irish potato famine pathogen Phytophthora infestans (Haas et al. 2009; Kroon et al. 2011; Yoshida et al. 2013). P. infestans, the causative agent of the late blight disease, is considered one of the most important biotic threats to global food security affecting worldwide production of key crops such as potato and tomato (Haverkort et al. 2008). Annual losses of potato harvest due to P. infestans would be sufficient to feed from 80 to many hundreds of millions of people and amount to several billion dollars per year (Cavalier-Smith 1999; Vleeshouwers et al. 2000; Win et al. 2012a).

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arms race with their host plants, and are continuously evolving new effector variants to evade NB-LRR immunity or suppress host defenses (Fry 2008; Win et al. 2012a).

To date, over 20 disease resistance (R) genes that function against *P. infestans* isolates have been cloned from various wild potato lines (*Solanum* spp.). All of these late blight R genes belong to the CC-NB-LRR class of NB-LRR receptor proteins and many occur in complex genetic loci (Jupe et al. 2012; Vleeshouwers et al. 2011). The *Solanum demissum* genes *R1*, *R2*, *R3*, *R4*, and *R10* have been bred into cultivated potato lines during the 20th century and have been widely used in conventional plant breeding programs (Vleeshouwers et al. 2011). However, these resistance genes have been rapidly defeated by new virulent races of *P. infestans* and are of limited value in modern agriculture (Fry and Goodwin 1997; Umaerus and Umaerus 1994; Wastie 1991).

One of the first late blight R genes to be bred in potato is *R3a*, which recognizes specific forms of *P. infestans* AVR3a, a host-translocated effector that belongs to the RXLR class of oomycete effectors (Armstrong et al. 2005; Birch et al. 2008; Huang et al. 2005; Morgan and Kamoun 2007). Two major *Avr3a* alleles occur in *P. infestans* populations. They encode *AVR3aK80/I103* (AVR3aKI) and *AVR3aE80/M103* (AVR3aEM), which differ in two amino acids in their effector domains. *R3a* responds to AVR3aKI but only weakly to AVR3aEM, and races of *P. infestans* that are virulent on *R3a* potato plants are homozygous for the AVR3aEM allele (Armstrong et al. 2005; Bos et al. 2006, 2009). AVR3aEM is the most prevalent allele in modern populations of *P. infestans* and the frequency of this allele has increased in the *P. infestans* population since the 19th century pandemic (Armstrong et al. 2005; Bos et al. 2009, 2010; Vleeshouwers et al. 2011; Yoshida et al. 2013). As a consequence, the *R3a* gene does not confer resistance to the majority of *P. infestans* strains that occur in modern agricultural systems.

The *P. infestans* AVR3a effector is a member of a large family of RXLR effectors that occur in most *Phytophthora* spp., including the pepper pathogen *P. capsici* and the soybean pathogen *P. sojae* (Bos et al. 2009; Haas et al. 2009; Tyler et al. 2006). Similar to other oomycete RXLR effectors, AVR3a proteins are modular, with the N-terminal region implicated in secretion and host translocation (Birch et al. 2008; Morgan and Kamoun 2007). The C-terminal domain carries the biochemical effector activity and is characterized by the WY α-helical fold that occurs in approximately 40% of *Phytophthora* RXLR effectors (Bouteny et al. 2011; Win et al. 2012b; Wirthenmueller et al. 2013). The majority of AVR3a family proteins, except for *P. sojae AVH1b*, do not elicit *R3a*-dependent hypersensitive cell death (Win et al. 2007). However, given their conservation across *Phytophthora* spp. and importance in pathogen virulence, AVR3a family effectors should be good targets for NB-LRR-mediated disease resistance. Indeed, *P. sojae AVR1b*, an AVR3a homolog, is known to mediate resistance in soybean plants carrying the NB-LRR gene *Rps1b* (Dou et al. 2008).

Although NB-LRR immune receptors have been known in plants for approximately 20 years, our mechanistic understanding of their function remains incomplete (Elmore et al. 2011; Maekawa et al. 2011b; Takken and Goverse 2012). According to current models of effector recognition, NB-LRR proteins can directly bind pathogen effectors or perceive them indirectly via effector-mediated modifications of plant proteins, that can be either guaerdes (vulurence targets of the effectors) or decoys (mimics of effector virulence targets) (Elmore et al. 2011; Takken and Goverse 2012; van der Hoorn and Kamoun 2008). The multidomain architecture of NB-LRR proteins enables them to operate as both pathogen sensors and activators of immune signaling (Takken and Goverse 2012). Intramolecular interactions between the different domains of NB-LRR proteins are important in maintaining them in an autoinhibited state that can be perturbed by pathogen effectors (Maekawa et al. 2011b; Takken and Goverse 2012). The NB domain, also known as NB-ARC (NB adaptor shared with APAF-1, plant resistance proteins, and CED-4), is thought to act as a molecular switch modulated by conformational changes and the replacement of ADP by ATP (Maekawa et al. 2011b; Takken and Goverse 2012). Recently, crystal structures of individual domains of plant NB-LRR proteins have begun to emerge (Bernoux et al. 2011; Hao et al. 2013; Maekawa et al. 2011a). Furthermore, the first structural insights into how different domains of NB-LRR proteins fold together have been obtained for the mouse protein NLRC4 but structural data of full-length plant NB-LRR receptors is still lacking (Hu et al. 2013).

A better understanding of NB-LRR protein structure and function would be useful to rationally design receptors with altered specificity. Previous studies have taken advantage of natural variation of R proteins and knowledge on NB-LRR structure to modify receptor specificity (Steinbrenner et al. 2012; Stirnweis et al. 2013). However, this strategy presents limitations and other methodologies are required when such natural variation is not available. An alternative approach is to screen randomly mutated NB-LRR proteins to recover variants with altered recognition properties. Such synthetic mutations may directly affect effector recognition specificity or result in sensitized (i.e., trigger-happy) NB-LRR receptors with a lower threshold for activation. Thus far, random mutagenesis has only been applied to the potato NB-LRR protein Rx in a domain-specific and stepwise manner (Farnham and Baulcombe 2006; Harris et al. 2013). Mutations in the LRR domain of Rx confer extended recognition against hitherto unrecognized strains of *Potato virus X* and *Poplar mosaic virus* (PopMV) (Farnham and Baulcombe 2006). Mutations that enhance activation sensitivity were recovered in a second round of mutagenesis, which focused on the N-terminal domain (Harris et al. 2013). These additional mutations near the NB pocket of Rx enhanced the response conferred by a single mutation in the LRR domain, resulting in resistance to PopMV (Harris et al. 2013).

In this study, we have extended the response spectrum of R3a to include the stealthy *P. infestans* AVR3a isoform AVR3aEM. To accomplish this, we screened a library of R3a variants obtained by random mutagenesis of the full-length R3a coding sequence, and recovered mutants that trigger hypersensitivity toward AVR3aEM while retaining recognition of AVR3aKI. We implemented the screen based on previous findings showing that the interaction between AVR3a and R3a can be reconstructed by concurrent expression in the model plant species *Nicotiana benthamiana* using *Agrobacterium tumefaciens* transient assays (agroinfiltration) (Armstrong et al. 2005; Bos et al. 2006). Single amino acid mutations that expand response to AVR3aEM were recovered in all three domains of R3a. One mutant R3aL668P, carrying a substitution in the N-terminal region of the LRR domain, responds to AVR3aEM more efficiently than the wild-type R3a. Remarkably, another R3a mutant, R3aN336Y, with an amino acid substitution near the NB site, gained the ability to respond to the *P. capsici* effector protein PcAVR3a. Our work contributes to understanding how NB-LRR receptor specificity can be modulated which, together with knowledge of pathogen effector diversity, can be exploited to develop synthetic immune receptors.

**RESULTS**

**R3a mutant library and screening strategy.**

To identify R3a mutants with expanded effector response, we first generated synthetic (non-natural) variants by polymerase chain reaction (PCR)-based random mutagenesis. We used
Fig. 1. R3a mutant library construction and screening strategy. The full length R3a coding sequence (3,849 bp) was used as template for a polymerase chain reaction-based random mutagenesis. Amplification products were cloned in the binary vector pCB302-3 (under transcriptional control of the Cauliflower mosaic virus 35S promoter) and transformed into Agrobacterium tumefaciens. The library was screened for gain-of-function phenotype by co-agroinfiltration in Nicotiana benthamiana of the R3a mutant clones (R3a*) with Phytophthora infestans AVR3aEM expressed from a Potato virus X (PVX)-based vector. Appearance of hypersensitive response (HR)-like phenotypes was evaluated at 5 days postinfiltration (dpi), and clones showing response to AVR3aEM were selected for a second round of co-infiltrations, including also co-expression with AVR3aKI and with a truncated version of green fluorescent protein (ΔGFP). Intensity of the observed responses was scored according to an arbitrary scale ranging from 0 (no HR-like phenotype) to 10 (confluent necrosis in the infiltrated area). Clones selected after these steps were named R3a+.
the full-length R3a coding sequence (3,849 bp) as template, and tested different conditions in order to obtain a mutation rate between one and two mutations every 1,000 bp. Mutagenized R3a molecules were cloned in the binary vector pCB302-3 (Xiang et al. 1999) under the transcriptional control of the Cauliflower mosaic virus 35S promoter, and transformed into A. tumefaciens to generate a library that could be screened by agroinfiltration assays in N. benthamiana. The pipeline to screen for R3a mutant clones with expanded response toward P. infestans AVR3aEM is illustrated in Figure 1. Briefly, we screened the mutant clones by co-expression with AVR3aEM in N. benthamiana leaves for gain of function assessment. We evaluated appearance of HR-like phenotypes at 5 days postinfiltration (dpi) and selected candidate clones for a second round of co-infiltrations, including co-expression with AVR3aKI to assess whether the R3a mutant clones retained their original activity, and with the negative control (truncated green fluorescent protein [ΔGFP]) to rule out that the observed cell death was a consequence of auto-activation. We scored the intensity of the observed responses according to an arbitrary scale ranging from 0 (no HR-like phenotype) to 10 (confluent necrosis in the infiltrated area). R3a mutant clones selected after implementation of this pipeline were designated R3a+.

R3a+: R3a mutants that respond to the AVR3aEM effector.

We screened approximately 2,200 clones of the R3a mutant library and identified 17 R3a+ clones that triggered visible HR in the presence of AVR3aEM (Fig. 2A and B). These candidate clones showed different degrees of response toward AVR3aEM. The R3a-17+, 18+, and 19+ mutant clones consistently gave the strongest HR with AVR3aEM. All 17 R3a+ clones retained the ability to respond to AVR3aKI (Fig. 2B). The majority of the R3a mutants did not trigger any HR symptoms when expressed with the negative control (ΔGFP). However, R3a clones 18+ and 19+ showed weak responses in some of the infiltrated leaves (HR index typically <2 on a scale of 1 to 10), indicating some degree of auto-activation (Fig. 2B). Overall, these results indicate that the response of R3a can be expanded toward P. infestans effector AVR3aEM without compromising its original recognition of AVR3aKI.

We next sequenced the 17 R3a+ clones to determine the underlying mutations. We found that each clone incorporated one to nine nonsynonymous mutations (on average approximately 4 amino-acid substitutions per R3a+ variant) (Fig. 2A). The mutations were distributed along the three NB-LRR protein domains (CC, NB-ARC, and LRR) (Supplementary Table S1). Interestingly, we noted a “hotspot” of amino acid changes around LRR repeats 15 and 18 (LRR repeat numbers throughout the article are based on Huang et al. 2005). This indicates that LRR 15 to 18 could be important for recognition or activation of R3a by AVR3aEM. Particularly, clone R3a-17+ had one single-amino-acid change in the LRR repeat 15 (K920 to E) compared with the wild-type R3a protein.

Single-amino-acid changes can expand R3a response to AVR3aEM.

To investigate which specific nonsynonymous mutations are responsible for the new response to AVR3aEM, we deconvoluted mutations observed in the original R3a+ clones into single-residue mutant clones. We co-infiltrated these single-residue

\[ \text{Fig. 2. Characterization of selected R3a mutant clones with expanded response to AVR3aEM (R3a+ clones).} \]

\[ \text{A, A total of 17 R3a+ clones were fully sequenced. Identified nonsynonymous mutations are indicated with stars positioned in the corresponding nucleotide-binding leucine-rich repeat-containing (NB-LRR) protein domain. Numbers of amino-acid changes related to wild-type R3a are indicated as bars on the left panel; a graph depicting number of amino-acid changes per LRR repeat for the 17 R3a+ clones is shown below.} \]

\[ \text{B, Hypersensitive response (HR) index scored at 7 days postinfiltration (dpi) after co-agroinfiltration in Nicotiana benthamiana of the R3a+ clones (under transcriptional control of the Cauliflower mosaic virus 35S promoter) or wild-type R3a with AVR3aEM, AVR3aKI, or a truncated version of green fluorescent protein (ΔGFP) expressed from a Potato virus X (PVX)-based vector. Bars represent the average of 10 replicas; the error bars represent standard deviation. Representative pictures for different values of HR index are included.} \]
mutant clones with AVR3aEM and scored HR-like phenotypes at 6 dpi. We observed that eight single-residue mutants triggered the HR in the presence of AVR3aEM with an HR index of 2 to 9, depending on the mutant (Fig. 3A). R3aC950R triggered the weakest response (HR index of approximately 2) whereas all the other mutants gave an HR-index ≥5 (Fig. 3A). All the tested clones retained the ability to trigger the HR when co-infiltrated with AVR3aKI (Fig. 3A). One of the mutant clones, R3aI148F, could also trigger the HR in the presence of the negative control ΔGFP (HR index of approximately 6), indicating that this mutation confers an auto-activation phenotype to R3a. Overall, these results indicate that single-amino-acid changes in R3a can expand its response to P. infestans AVR3aEM.

To date, the three-dimensional structure of a full-length plant NB-LRR protein has not been determined. However, representative structures of each domain (CC, NB-ARC, and LRR) are available from either plant or other eukaryotic proteins. In an attempt to understand the structural consequences of the R3a+ single mutations, we generated homology models of the individual CC, NB-ARC, and LRR regions of R3a using protein fold-recognition algorithms, as implemented in Phyre2 (Kelley and Sternberg 2009). Of the eight single-residue mutants described above, six result in amino acid substitutions in the LRR domain (R3aL668P, R3aK920E, R3aE941K, R3aC950R, R3aE983K, and R3aK1250R), one in the CC domain (R3aI148F), and one in the NB-ARC domain (R3aN336Y). We mapped the position of residues contained within the NB-ARC and LRR domains on the respective homology models we generated (Fig. 3B). Unfortunately, although we were able to generate a model for the R3a CC domain, it did not include the region containing the I148F mutant; therefore, it is not presented here. Such homology models can, typically, only be indicative of the position of individual amino acids. However, it is noteworthy that R3aN336Y maps near the NB site, at the junction of the NB and ARC1 regions, and close to the P-loop, sensor 1, GLPL, and MHD motifs that have previously been implicated in molecular switch events associated with nucleotide exchange in plant NB-LRR proteins (van Ooijen et al. 2008). The N336Y mutation places a bulky hydrophobic amino acid in place of a smaller residue and the resulting changes in the structure, due to steric constraints, may explain the observed phenotype. Although not included in a homology model, the effects of the I148F mutant may also be explained through the substitution of a larger residue in the place of a smaller one. The remaining six mutations (R3aL668P, R3aK920E, R3aE941K, R3aC950R, R3aE983K, and R3aK1250R) are located in the LRR domain. Four of these mutations are clustered around LRR repeats 15 and 18 (Huang et al. 2005). The other mutations are distributed along and around the solenoid. Some of the mutations in the LRR domain are predicted to be in residues exposed on the protein surface and others buried in the hydrophobic core of the repeats.

The R3aL668P mutant shows increased response to AVR3aEM compared with R3a.

To determine whether the R3a+ single-residue mutants have altered sensitivity to AVR3aKI compared with wild-type R3a, we co-infiltrated (side by side) N. benthamiana leaves with A. tumefaciens strains expressing mutant and wild-type R3a constructs mixed with different dilutions of A. tumefaciens strains expressing AVR3aKI. We then scored each pair for the combination giving the strongest HR (Fig. 4A; Supplementary Fig. S1). R3aL668P consistently (>60% of infiltration sites) triggered a stronger HR than R3a when co-infiltrated with AVR3aKI whereas R3aC950R, R3aE983K, and R3aK1250R occasionally (approximately 10% of infiltrations) triggered a stronger HR with AVR3aKI. We further confirmed that R3aL668P has an enhanced response to AVR3aKI independently of the dilution level of the A. tumefaciens strain carrying the effector (Fig. 4B). These findings indicate that this single-amino-acid change in R3a (L668P) not only expands the response of R3a to AVR3aEM but also increases the sensitivity of R3a for its cognate effector AVR3aKl.

R3a+ single-residue mutants respond to multiple variants of AVR3a.

Although all the R3a+ mutants we identified have expanded response to AVR3aEM, they may not have the same response spectrum to additional variants of AVR3a. To test whether the activity acquired by R3a+ resistance proteins extends to additional variants of AVR3a besides AVR3aKl and AVR3aEM, we co-infiltrated the R3a+ single-residue mutants with a set of previously described AVR3a mutants (Bos et al. 2009). We selected AVR3a variants with Pro, Asp, Tyr, Phe, or Trp residues at position 80 (AVR3aE80K/M103) because they do not trigger the HR when co-expressed with wild-type R3a in a wound-inoculation assay (Bos et al. 2009). Interestingly, the R3a+ single-residue mutants showed variable response spectra to these AVR3a variants (Fig. 5; Supplementary Fig. S2). R3aE983K showed the widest spectrum, responding with an HR index of >7.5 to all tested AVR3a variants, followed by clones R3aL668P, R3aK920E, R3aN336Y, and R3aE983K (Fig. 5B). These results indicate that single-amino-acid changes in R3a can dramatically expand the response of this NB-LRR protein not only to AVR3aEM but also to multiple artificial variants of AVR3a. The findings also show that the R3a mutants we recovered have variable response activities to AVR3a mutants.

The R3aN336Y mutant can respond to PcAVR3a4, an AVR3a homolog from P. capsici.

We hypothesized that some of the R3a+ mutants we identified may respond to members of the AVR3a family from another Phytophthora sp. To test this, we co-expressed the eight single-residue R3a+ mutants with several AVR3a-like effectors from the pepper and cucurbit pathogen P. capsici (Bos 2007). Remarkably, R3aN336Y, which carries an amino-acid substitution near the predicted NB pocket of R3a (Fig. 3B), consistently responded to the effector PcAVR3a4 (HR index approximately 6) whereas the wild-type R3a did not (HR index = 0) (Fig. 6). R3aI148F, which is mutated in the CC domain and was previously tagged as an autoactive mutant (Fig. 3A), also responded to PcAVR3a4 (Fig. 6A). These results indicate that the single-amino-acid substitution N336Y can expand R3a response not only to AVR3aEM but also to another AVR3a family member from P. capsici.

DISCUSSION

The potato NB-LRR immune receptor R3a mediates resistance to strains of P. infestans that secrete the AVR3aKl effector but not to strains that only produce the AVR3aEM isoform. AVR3aEM is the product of an AVR3a allele that is predominant in modern populations of P. infestans, meaning that R3a is a “defeated” disease resistance gene in most potato production areas (Armstrong et al. 2005; Bos et al. 2009, 2010; Yoshida et al. 2013). In this study, we took the first steps to resurrect R3a by successfully screening and identifying R3a+ mutants that respond to both AVR3aKI and AVR3aEM (Figs. 1 and 2). In total, we identified eight single-amino-acid mutations that expanded R3a response to AVR3aEM without compromising its activity toward AVR3aKI (Fig. 3). The mutants span the length of the R3a protein, providing unbiased and valuable insight into structure–function relationships in this immune receptor. Remarkably, one of the R3a mutants, R3aN336Y, gained the capacity to respond to an AVR3a homolog from another oomycete.
Fig. 3. Single amino acid changes located at different nucleotide-binding leucine-rich repeat-containing (NB-LRR) domains can expand R3a response to AVR3aEM. A, Single-residue mutation clones derived from the original R3a+ and wild-type R3a (all under transcriptional control of the Rpi-vnt1.1 promoter) were co-agroinfiltrated in Nicotiana benthamiana with AVR3aEM, AVR3aKI, or a truncated version of green fluorescent protein (ΔGFP) expressed from a Potato virus X (PVX)-based vector. The empty vector (E.V.) clone was included as control. Hypersensitive response index scored at 6 days postinfiltration (dpi) is plotted; bars represent the average of 13 to 18 replicas for ΔGFP and AVR3aKI and 28 to 30 replicas for AVR3aEM; error bars represent standard deviation. B, Schematic (cartoon) representation of the homology models generated for the NB-ARC and LRR regions of the R3a protein. Models were generated using the Phyre2 server (Kelley and Sternberg 2009). The NB-ARC is shown in purple and the LRR in orange. The carbon atoms of the single-residue R3a+ mutations are shown in red and the carbon atoms of ATP are shown in yellow. The inset shows a close-up of the ADP/ATP binding site within the NB-ARC domain (here, carbon atoms are colored in cyan [P-loop], pink [sensor 1], green [GLPL motif], and orange [MHD motif]). Mutation K1250R is not shown because the region containing this residue was not modeled.
species, the vegetable pathogen *P. capsici*. These results validate the strategy of Farnham and Baulcombe (2006) and Harris and colleagues (2013) in demonstrating that synthetic alleles of disease resistance genes with expanded specificity spectrum can be produced in the laboratory.

We screened a library of approximately 2,200 clones with AVR3aEM and recovered 17 R3a+ clones (Figs. 1 and 2). This represents an approximately 0.8% success rate. Farnham and Baulcombe (2006) achieved a similar success rate because they recovered 13 gain-of-function mutants of Rx after screening 1,920 mutants (approximately 0.7%). Given that our mutants have an average rate of nonsynonymous mutations of 4 and that R3a is 1,282 amino acids in length, the mutant library we generated had approximately 7x coverage for each amino acid (2,200 × 4/1,282 = 6.86). The depth of the screen is validated by the identification of a cluster of four mutations around LRR repeats 15 and 18 (Fig. 3B). However, because most mutant clones have multiple amino acid substitutions, we may have failed to recover mutants with compensating beneficial and detrimental mutations. Therefore, the screen, although successful, may have missed additional useful mutations. Nonetheless, the 7x deep R3a mutant library we constructed is a permanent resource that can be rescreened with additional forms of AVR3a from either *P. infestans* or other *Phytophthora* spp.

Two general mechanisms could underlie the ability of the R3a+ mutants to respond to additional *Phytophthora* effectors,

![Graph showing percentage of infiltrated sites](image)

**Fig. 4.** The R3aL668P mutant shows increased response to AVR3aK1 compared with R3a. **A,** The single-residue mutant R3a+ clones or wild-type R3a (all under transcriptional control of the *Rpi-vnt1.1* promoter; optical density at 600 nm [OD$_{600}$] = 0.3) were co-agroinfiltrated side by side with serial dilutions of AVR3aK1 (under transcriptional control of the Cauliflower mosaic virus 35S promoter) in *Nicotiana benthamiana* (from the tip to the base of the leaf, expressed in OD$_{600}$ = 0.04 AVR3aK1 + 0.11 empty vector [E.V.], 0.075 AVR3aK1 + 0.075 E.V., and 0.15 AVR3aK1). The E.V. clone was included as control. Hypersensitive response was scored at 4 days postinfiltration (dpi) for neighbor spots (R3a+ and wild-type R3a in opposite sides of the leaf) as follows: “R3a+ stronger than R3a”, “R3a+ equal to R3a”, or “R3a+ weaker than R3a”. Results for the lowest concentration of AVR3aK1 are plotted as percentage of compared infiltrated areas showing each of the possible outcomes. **B,** A representative picture of an *N. benthamiana* leaf co-agroinfiltrated as explained in A with the R3a+ clone R3aL668P and serial dilutions (upper = higher, inferior = lower) of *Agrobacterium tumefaciens* cells carrying the AVR3aK1 construct (as detailed above). The picture was taken at 4 dpi.
Fig. 5. R3a+ mutants respond to multiple AVR3a variants. A, Schematic representation of Phytophthora infestans effector protein AVR3a, showing polymorphic amino acid positions 80 and 103 in the effector domain. B, R3a+ single-residue mutants and wild-type R3a (all under transcriptional control of the Rpi-vnt1.1 promoter) were co-agroinfiltrated in Nicotiana benthamiana with a truncated version of green fluorescent protein (ΔGFP), AVR3aK1, AVR3aEM, and a collection of AVR3a mutants with amino-acid substitutions (Pro, Asp, Tyr, Phe, or Trp) at position 80 in AVR3aE80X/M103. All these AVR3a variants were expressed from a Potato virus X (PVX)-based vector. The empty vector (EV) clone was included as control. Hypersensitive response (HR) index scored at 5 days postinfiltration (dpi) is plotted; columns represent the average of at least 12 replicas and error bars represent standard deviation.
The R3a+ mutants could be altered in recognition specificity, gaining the ability to sense a wider spectrum of effectors, either directly or indirectly. Alternatively, the R3a+ mutants could have increased activation sensitivity. Sensitized or “trigger-happy” mutants would have a lower threshold for activation, producing a stronger response to weak elicitors such as AVR3aEM and PcAVR3a4 than wild-type R3a protein. According to current knowledge on structure–function relationships of NB-LRR proteins, mutations in the C-terminal half of the LRR domain are expected to affect recognition specificity, whereas those located in the CC, NB-ARC, and N-terminal region of the LRR domain are more likely to affect activation and signaling threshold (Takken and Goverse 2012). Therefore, the eight R3a+ amino-acid mutations may affect different aspects of the response to AVR3a.

**Fig. 6.** The R3aN336Y mutant responds to PcAVR3a4, an AVR3a homolog from the vegetable pathogen *Phytophthora capsici*. A, The R3a+ single-residue mutants and wild-type R3a (all under transcriptional control of the *Rpi-vnt1.1* promoter) were co-agroinfiltrated in *Nicotiana benthamiana* with a truncated version of green fluorescent protein (ΔGFP), AVR3aKI, AVR3aEM, and the AVR3a family member from *P. capsici*, PcAVR3a4. All these effectors were expressed from a *Potato virus X* (PVX)-based vector. The empty vector (E.V.) clone was included as control. Hypersensitive response index scored at 5 days postinfiltration (dpi) is plotted; columns represent the average of at least 10 replicas and error bars represent standard deviation. B, A representative picture of an *N. benthamiana* leaf co-agroinfiltrated as explained in A, with PcAVR3a4 in combination with E.V., wild-type R3a, or the R3a+ clones R3aI148F and R3aN336Y. The picture was taken at 5 dpi.
effectors given that they are spread along the R3a protein (Fig. 3), with R3aK1250R, R3aK366Y, and R3aL668P corresponding to regions possibly involved in signal transduction and R3aK320E, R3aE941K, R3aC950R, R3aE983K, and R3aK1250R located in the region of the protein that might influence specificity.

Four of the eight amino-acid substitutions that enhance the response to AVR3aEM map to the central region of the LRR domain (repeats 15 to 18) (Fig. 3), pointing to a potential hot-spot. Of these mutations, C950R is phenotypically distinct, conferring a relatively weak response to AVR3aEM and other AVR3a variants (Figs. 3A and 5B). The three remaining mutants (K920E, E941K, and E983K) are phenotypically similar in all the assays we tested. All three mutants carry substitutions in charged amino acids that resulted in a switch from a positively to a negatively charged residue, or vice-versa. These changes are consistent with an altered binding state of the R3a protein to a ligand that could be either the effector or a plant protein. Interestingly, LRR 15 to 18 exactly match the region identified by van Ooijen and colleagues (2010) as the binding site of I-2, an R3a homolog from tomato, to the small heat shock protein 20 RS12. These authors showed that RS12 is required for I-2 function, possibly by affecting stabilization of this NB-LRR protein. Therefore, mutations in this central region of the LRR domain could alter binding to a plant chaperone protein such as RS12, therefore perturbing steady-state levels of R3a. Alternatively, the central region of the LRR domain could be the binding site to the AVR3a effector or a protein complex between the effector and host proteins.

The L668P mutation in the fifth LRR is the only one of the eight mutations that resulted in significantly increased response to AVR3aEM, and appears to act by sensitizing the R3a protein (Fig. 4). This finding confirms previous observations implicating the N terminus of the LRR domain in signalizing (Bendahmane et al. 2002; Farnham and Baulcombe 2006; Takken and Goverse 2012). The N-terminal region of the LRR domain is thought to fold onto the NB-ARC domain to stabilize the NB-LRR protein, maintaining it in an autoinhibited state (Takken and Goverse 2012). The Pro at position 668 may perturb the conformation of the N terminus of the LRR domain, resulting in altered intramolecular interaction with the NB-ARC domain. Importantly, the R3aL668P mutant is not autoactive (Figs. 3A, 5B, and 6A). Therefore, mutants in the N terminus of the LRR domain can be used to optimize NB-LRR protein activity and modulate the response to both weakly and strongly recognized effectors. Such mutants could be useful for enhancing the response of an NB-LRR receptor to a pathogen effector without the negative effects of effector-independent activation.

The NB-ARC domain mutation N336Y was the only one that resulted in a gain of response to an AVR3a-like effector from a different Phytophthora sp., without triggering autoactivity (Fig. 6). This finding is reminiscent of the results of Farnham and Baulcombe (2006), which showed that single-residue LRR mutants of the Rx protein (mutants M1 to M4) respond to the coat protein (CP) of a new virus, PopMV. Farnham and Baulcombe (2006) concluded that Rx mutants M1 to M4 gained recognition of the PopMV CP. Subsequently, Harris and colleagues (2013) showed that the RxM1 mutant can be enhanced by sensitizing mutations in the NB-ARC domain, and that without M1 these sensitized mutants (RsX1 to RsX54) do not produce a robust response to PopMV CP. In our case, a single mutation in the NB-ARC domain of R3a was sufficient to confer a substantial response to the P. capsici effector PcAVR3a4 (Fig. 6). Although, phenotypically, the R3aN336Y mutant seems to be similar to the M1 to M4 mutations, it seems unlikely that a mutation near the NB site would affect recognition per se. Perhaps R3aN336Y is an oversensitized mutant that reveals cryptic activities of R3a. If correct, this model implies that immune receptors like R3a have a much greater capacity to recognize diverse pathogen effectors than previously thought. Mutations similar to N336Y have the potential to reveal cryptic activities of immune receptors with potentially important implications for agriculture. In addition, the unexpected gain-of-function conferred by the N336Y mutation highlights the value of mutagenizing full-length NB-LRR proteins without preconceived assumptions about the LRR being a recognition domain (Farnham and Baulcombe 2006).

This work has implications for crop improvements. However, preliminary infection assays, conducted after transient and stable expression of the R3a+ mutants described here, failed to detect increased resistance to P. infestans in either N. benthamiana or potato plants. The response of the R3a+ clones to AVR3aEM is variable in strength (Fig. 3A) but, in general, it remains weaker than the response to AVR3aK1. Assaying different residues in each of the R3a+ mutant positions or combining mutations in the same R3a+ clone could help to optimize the strength of the response to AVR3aEM. In particular, this last approach proved to be valid for Rx NB-LRR, because combination of mutations in different domains led to a more robust response (Harris et al. 2013). Combination of two or more single-amino-acid changes identified in R3a+ clones may be required to assemble a more stable recognition complex in order to elicit resistance. Screening for the appearance of the HR has proven to be successful in selecting for NB-LRR clones that confer resistance to pathogens (Farnham and Baulcombe 2006). Nevertheless, in some instances, host-programmed cell death can be uncoupled from resistance (Century et al. 1995; Hao et al. 2013; Heidrich et al. 2012), and we cannot ignore the possibility that the cell death response triggered by the R3a+ mutants in the presence of AVR3aEM may take place in an inadequate spatiotemporal frame that compromises resistance. In addition, it is unclear whether the recognition of AVR3a by R3a is direct or dependent on host-specific proteins (guardce, decay, or co-factor proteins) (Bos et al. 2009; Engelhardt et al. 2012). To date, yeast two-hybrid and co-immunoprecipitation experiments have failed to show a direct interaction between AVR3a and R3a (Engelhardt et al. 2012). Clearly, a better understanding of how R3a functions is required to translate the gain of effector response noted in the R3a+ mutants into plant disease resistance.

The ability to extend the response of an NB-LRR protein to a wider range of related effectors offers the opportunity to develop broad-spectrum solutions to plant pathogens. In the case of the AVR3a family, multiple members exist in phylogenetically diverse Phytophthora spp. (Haas et al. 2009; Tyler et al. 2006). As Phytophthora genome sequences become available, it will be possible to identify AVR3a-like proteins and screen them with the R3a mutant library. Artificial evolution by random mutagenesis complements site-directed mutagenesis and domain shuffling approaches to improve NB-LRR proteins (Steinbrenner et al. 2012; Stirnweis et al. 2013; Takken and Goverse 2012). In the future, combining these different approaches could be used to generate synthetic R genes with novel beneficial properties. In addition, with the advent of plant genome editing, such beneficial mutations may be introduced into an elite crop cultivar genome without the insertion of additional sequences (Belhaj et al. 2013; Li et al. 2013; Mussolin and Cathomen 2013; Nekrasov et al. 2013; Schornack et al. 2013; Shan et al. 2013).

MATERIALS AND METHODS

Microbial strains and growth conditions.

A. tumefaciens GV3101 (Hellens et al. 2000) was used in molecular cloning experiments and library construction, and was
R3a mutant library.

Random mutagenesis of the full-length R3a coding sequence was performed using the Diversify PCR random mutagenesis kit from Clontech (Takara Bio Company, Mountain View, CA, U.S.A.). PCR amplification was performed using S. demissum R3a gene as template (Huang et al. 2005), following the provider’s protocol. Primers were designed to amplify the R3a coding sequence, including restriction enzymes recognition sites for subcloning (R3a_BamHI_FW: 5’-GGAGGATCCCAT GGGATTGCTTACGC-3’ and R3a_SpeI_Rev: 5’-GGA ACTAGTTCACATGCATTCC-3’). Different conditions were tested to generate various mutation rates. Conditions that gave a mutation rate of one to two mutations every 1,000 bp (four to eight in a 4,000-bp gene) were selected. The PCR reaction was purified using a QIAquick PCR purification kit (Qiagen, Hilden, Germany). The purified PCR product and the binary vector pCB302-3 (Xiang et al. 1999) were digested with BamHI and SpeI restriction enzymes (Roche Applied Science, Penzberg, Germany). The digested PCR products and vector were gel-purified using the kit Wizard SV Gel and PCR clean-up system (Promega Corp., Madison, WI, U.S.A.), and ligation mixes were directly transformed into A. tumefaciens GV3101 by electroporation. Transformants were picked into 384-well plates using a QPiX colony-picking robot (Genetix, New Milton, U.K.) for preparation of freeze stocks. The library was kept at −80°C.

R3a mutant library screening.

The screening was performed by agroinfiltration transient assays in N. benthamiana (Bos et al. 2006, 2009; Van der Hoorn et al. 2000). In all, 2.0 ml 96 Deepwell plates (StarLab, Hamburg, Germany) containing 500 µL of LB media with antibiotics (rifampicin at 50 µg/liter, gentamicin at 20 µg/liter, and kanamycin at 50 µg/liter) were inoculated with the library clones and grew at 28°C for 48 h (to reach an optical density at 600 nm [OD600] of 1 to 1.2). Cultures were pelleted by centrifugation (5 min at 3,500 rpm and 15°C) and resuspended with infiltration buffer (0.5% MS salts, 10 mM MES, 2% sucrose, and 200 µM acetylsyringone, pH 5.6) to a final OD600 of 0.6. A. tumefaciens GV3101 transformed with pGR106-FLAG-AVR3aΔK80I103 (AVR3aKI) and pGR106-FLAG-AVR3aE103ΔK80I103 (AVR3aEM) (Armstrong et al. 2005; Bos et al. 2006, 2009) were grown for agroinfiltration as previously described (Van der Hoorn et al. 2000), except that LB media supplemented with rifampicin at 50 µg/liter, gentamicin at 20 µg/liter, and kanamycin at 50 µg/liter was used. Cultures were pelleted and resuspended in infiltration buffer as described above. For transient co-expression of R3a, R3a mutant clones, and A VR3a, the cells resuspended in infiltration buffer were mixed to have a final OD600 = 0.3 (R3a clones) and 0.15 (AVR3a clones). A. tumefaciens GV3101 transformed with pGR106-AGFP (containing a truncated version of gfp as described by Bos and colleagues [2006]) was grown as the AVR3a clones. Agroinfiltration experiments were performed on 4-week-old N. benthamiana plants. Plants were grown and maintained throughout the experiments in a controlled environment room with a temperature of 22 to 25°C and high light intensity. HR phenotype development was monitored from 3 to 8 dpi.

Agroinfiltration assays.

Selected R3a+ clones were co-agroinfiltrated in N. benthamiana leaves to compare their relative response to different effec-
BamHI-Fw (S′-GGAGGATCCCAATGGAGATTGCTTAG CAGTGTGTTG-3′) and PCB302-3′rv (S′-ACATGCTTTACGCT AATTCACACAG-3′). The PCR amplification product was digested with BamHI and XhoI restriction enzymes, gel purified, and ligated into pCBNpTi-1-vnt1.1/PT previously treated with the same restriction enzymes.

**R3a structure predictions.**

Homology models of the individual CC, NB-ARC, and LRR regions of R3a were generated using protein fold recognition algorithms, as implemented in Phyre2 (Kelley and Sternberg 2009), and sequences covering the individual domains. Phyre2 revealed the highest score for the CC domain using the CC region of MLA10 as a template (PDB file 3QFL, confidence 99.7%, sequence identity 15%). However, this model only covered the region between amino acid 14 and 131 and did not show the position of 1148. Searches with the sequence of the NB-ARC region showed the highest modeling score using CED4 as a template, and this generated a model spanning residues 181 to 515 (PDB file 2ASY, confidence 100%, sequence identity 16%). The template used for the LRR region was the toll-like receptor 3 ligand-binding domain (PDB file 2AOZ, confidence 100%, sequence identity 14%). Because the resulting model covers residues 584 to 1,206, mutation K1250R is not shown in Figure 3B. Note that the LRR repeat numbering is based on Huang and colleagues (2005).

**ACKNOWLEDGMENTS**

This work was supported by the European Research Council (ERC, proposal NGRB), the U.K. Biotechnology and Biological Sciences Research Council (BBSRC grants BB/H019813/1, the Gatsby Charitable Foundation, and the John Innes Foundation. M. E. Segretin was supported by the Consejo Nacional de Investigaciones Científicas y Técnicas (CONICET). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript. We thank A. Giannakopoulou, S. Schornack, N. Chambouret, B. Staskawicz, E. Huijtema, S. Huang, V. Vleeshouwers, I. Hein, and P. Birch for useful discussions or for providing biomaterial; K. Kowitwanich for technical support; G. Bennett for technical advice; and A. Bennaceur for promoting our collaborative interactions. M. E. Segretin, M. Pais, M. Franceschetti, A. Chaparro-Garcia, M. J. Banfield and S. Kamoun conceived and designed the experiments. M. E. Segretin, M. Pais, M. Franceschetti, A. Chaparro-Garcia, and J. I. Bos performed the experiments. M. E. Segretin, M. Pais, M. Franceschetti, M. J. Banfield and S. Kamoun analyzed the data. M. E. Segretin, M. Pais, M. Franceschetti, M. J. Banfield and S. Kamoun wrote the paper.

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Food and Agriculture Organization of the United Nations (FAO) website: faostat.fao.org/site/339/default.aspx