





Multiple pairs of allelic MLA immune receptor-powdery mildew AVR_A effectors argue for a direct recognition mechanism

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Abstract Nucleotide-binding domain and leucine-rich repeat (NLR)-containing proteins in plants and animals mediate intracellular pathogen sensing. Plant NLRs typically detect strain-specific pathogen effectors and trigger immune responses often linked to localized host cell death. The barley *Mla* disease resistance locus has undergone extensive functional diversification in the host population and encodes numerous allelic NLRs each detecting a matching isolate-specific avirulence effector (AVR_A) of the fungal pathogen *Blumeria graminis* f. sp. *hordei* (*Bgh*). We report here the isolation of *Bgh* AVR_{a7}, AVR_{a9}, AVR_{a10}, and AVR_{a22}, which encode small secreted proteins recognized by allelic MLA7, MLA9, MLA10, and MLA22 receptors, respectively. These effectors are sequence-unrelated, except for allelic AVR_{a10} and AVR_{a22} that are co-maintained in pathogen populations in the form of a balanced polymorphism. Contrary to numerous examples of indirect recognition of bacterial effectors by plant NLRs, co-expression experiments with matching *Mla-AVR*_a pairs indicate direct detection of the sequence-unrelated fungal effectors by MLA receptors. DOI: https://doi.org/10.7554/eLife.44471.001

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Introduction

The NLR family of immune receptors is structurally conserved between animals and plants, perceives non-self and modified-self molecules inside host cells, and mounts potent innate immune responses to terminate microbial pathogenesis (*Maekawa et al., 2011a*). Animal NLRs are normally activated by conserved microbe- or damage-associated molecular patterns (MAMPs/DAMPs), whereas plant NLRs typically detect strain-specific pathogen effectors, designated avirulence effector proteins (AVRs) (*Maekawa et al., 2011a; Jones et al., 2016*). Plant NLRs recognize either the effector structure or sense effector-mediated modifications of host proteins (*Jones et al., 2016*). Single plant NLRs or NLR pairs can also detect the manipulation of effector target mimics, called decoys, and a variation of the latter mechanism involves the direct integration of decoy domains mimicking host targets within NLRs, termed integrated decoy domains (*Jones et al., 2016*; *Cesari et al., 2014*; *Kroj et al., 2016*). The selective forces shaping the evolution of the comparatively small complement of NLRs in vertebrates are incompletely understood (~20 family members, (*Lange et al., 2011*; *Meunier and Broz, 2017*)), but in plant species co-evolution with host-adapted pathogens has strongly influenced the expansion and diversification of NLR repertoires (*Jacob et al., 2013*; *Meyers et al., 2005*).



eLife digest Powdery mildews are fungal diseases that affect many plants, including important crops such as barley. The fungi behind these diseases deliver molecules known as effectors inside plant cells, which manipulate the plants' biology and help the fungus to invade the plants' tissues. In response, some plants have evolved immune receptors encoded by so-called *R* genes (short for resistance genes) that detect the effectors inside the plant cell and trigger an immune response. The response often kills the plant cell and those nearby to limit the spread of the fungus. Effectors that are recognized by host immune receptors are termed avirulence effectors (or AVRs for short).

Scientists tend to assume that most effectors do not bind directly to their immune receptors. Instead, it is thought that the immune receptors are more likely to be detecting a change in some other plant protein that is caused by the effectors' activities.

In barley populations, one *R* gene that protects against powdery mildew encodes an immune receptor known as MLA. Different plants can carry subtly different versions of this *R* gene meaning that they make similar but different variants of the same receptor. Each MLA variant confers immunity only to strains of powdery mildew that carry the matching AVR effector. A few AVR effectors from powdery mildews have been identified, but most AVR effectors from powdery mildews remain unknown.

Saur et al. looked for new AVR effectors from powdery mildew fungi collected in the field, and found four that were recognized by barley plants carrying MLA variants. Two of these new effectors were fairly similar to each other, but they were all unlike those that had been identified previously.

When Saur et al. engineered barley cells to make these new AVRs alongside their matching MLA receptors, the cells died – which is consistent with the expected immune response. Similar experiments with distantly related tobacco plants agave the same results. This suggested that the immune receptors did not need any other barley proteins to recognize the effectors, indicating that the interaction between the two may be direct. Indeed, two other techniques that test for direct protein-protein interactions, – one that involved extracts from tobacco leaves, and another that involved yeast, – gave results consistent with a direct interaction between the MLA receptor variants and the fungal effectors.

Plant disease is still a major cause of loss of yield in crops. Transferring an R gene from one plant species to another is a potentially powerful approach to help crops resist disease. The discovery that multiple variants of the same resistance gene can bind to dissimilar effectors from a disease-causing fungus in distantly related plant species underlines the potential of this approach.

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Plant disease resistance (R) genes to host-adapted pathogens often encode NLRs, are frequently members of large gene families organized in complex clusters of paralogous genes, and can rapidly evolve through a range of natural gene diversification mechanisms (Jacob et al., 2013; Meyers et al., 2005). There are several examples of allelic series of NLR-type R genes known in plants (Ellis et al., 1999; Allen et al., 2004; Srichumpa et al., 2005; Seeholzer et al., 2010; Kanzaki et al., 2012). In these cases, multiple distinct recognition specificities evolved in the host population at a single R gene with each allele detecting a corresponding strain-specific AVR in the pathogen population. Such multi-allelic NLR-type R genes are particularly instructive for studying the underlying co-evolutionary process between host and pathogen.

Ascomycete powdery mildews are widespread pathogens of thousands of angiosperm plant species in temperate climates, including economically relevant crops (*Glawe*, 2008). They are obligate biotrophic pathogens, meaning that their growth and reproduction is entirely dependent on living host cells. The haploid barley powdery mildew pathogen *Blumeria graminis* forma specialis *hordei* (*Bgh*) multiplies mainly clonally and is a member of the species *Blumeria graminis* that is specialized for its host plant barley (*Hordeum vulgare*). There are various specialized forms (*formae speciales* or fs. spp.) of *B. graminis*, each of which is capable of infecting the respective host plant species belonging to the grass (Poaceae) family, including cereals such as barley and wheat (*Wyand and Brown*, 2003). Within each *forma specialis*, numerous isolates (strains) can be distinguished in the pathogen population, based on their respective virulence or avirulence infection phenotypes vis-à-



vis particular genotypes of the host population (*Lu et al., 2016*). The genomes of powdery mildews are characterized by the loss of several, otherwise widely conserved Ascomycete genes with functions related to carbohydrate degradation and primary and secondary metabolism (*Spanu et al., 2010*; *Wicker et al., 2013*), and this is believed to explain their strict dependence on living plant cells. Similar to other filamentous phytopathogens, grass-infecting powdery mildew genomes harbor hundreds of candidate secreted effector protein (CSEP)-coding genes, which are assumed to contribute to fungal pathogenesis (*Wicker et al., 2013*; *Pedersen et al., 2012*; *Guttman et al., 2014*). Pathogen effectors often work by subverting innate immune responses, thereby facilitating host colonization and disease (*Rovenich et al., 2014*).

Domesticated barley and wheat contain numerous powdery mildew R gene loci that were often introgressed from their corresponding wild relatives (Jørgensen and Wolfe, 1994; Lutz et al., 1995; Maekawa et al., 2019). In both barley and its close relative wheat, one of these powdery mildew R loci, designated mildew locus a (Mla) and powdery mildew 3 (Pm3), respectively, has been subject to exceptional functional diversification, resulting in large numbers of Mla or Pm3 recognition specificities (Seeholzer et al., 2010; Lutz et al., 1995; Bhullar et al., 2010). Although wheat Pm3 and barley Mla loci each span a cluster of NLR genes, known Pm3 and Mla recognition specificities to the B. graminis pathogen appear to have arisen from allelic diversification of a single NLR gene in the corresponding NLR clusters (Seeholzer et al., 2010; Maekawa et al., 2019; Shen, 2003). Isolate-specific disease resistance to Bgh mediated by MLA receptors is invariably linked to the activation of localized host cell death, and this immune response likely terminates growth of the biotrophic pathogen by shutting off its nutrient supply (Boyd et al., 1995). Of note, the Mla orthologs Sr33 in wheat (Periyannan et al., 2013) and Sr50 in rye (Mago et al., 2015) confer disease resistance to the stem rust pathogen Puccinia graminis f. sp. tritici (Pgt) isolate Ug99, a major threat to global wheat production. Pgt and Bgh belong to the Basidiomycota and Ascomycota phyla, respectively, indicating that MLA receptors can detect the presence of independently evolved avirulence effectors. Barley Mla and wheat Pm3 both encode intracellular NLRs with an N-terminal coiled-coil (CC) domain but lack significant sequence relatedness (Zhou et al., 2001), whereas 23 allelic barley MLA resistance proteins exhibit >91% amino acid (aa) sequence identity (Seeholzer et al., 2010), and 17 deduced allelic wheat Pm3 receptors share >97% as sequence identity (Bhullar et al., 2010). Diversifying selection among resistance alleles of Mla and Pm3 is largely confined to regions encoding the C-terminal leucine-rich repeats (LRR) of the receptors (Seeholzer et al., 2010; Maekawa et al., 2019; Bhullar et al., 2010). The polymorphic MLA LRR is critical for effector detection as shown by a series of reciprocal domain swaps between MLA1 and MLA6 (Shen, 2003). Barley Mla1 confers race-specific disease resistance to a Bgh isolate carrying the cognate avirulence gene AVR_{a1} in transgenic Arabidopsis thaliana, suggesting ~ 150 million years of evolutionary conservation of the underlying immune mechanism and potentially pointing to a direct recognition mechanism of AVR_A effectors that is highly conserved between barley and A. thaliana (Maekawa et al., 2012).

Recently, the *Bgt* avirulence gene *AvrPm3*^{a2/f2}, recognized by the wheat *Pm3a* and *Pm3f* alleles, was identified by a map-based cloning approach and found to encode a typical CSEP that belongs to an effector family of 24 members (*Bourras et al., 2015*). The clonal nature of the haploid *Bgh* pathogen facilitated the identification of isolate-specific sequence variation in the transcriptomes of a global collection of 17 *Bgh* strains. These were associated with *Bgh* pathotypes on *Mla1-* and *Mla13*-containing near-isogenic barley lines to identify sequence-unrelated *AVR_{a1-}* and *AVR_{a13-}* encoding CSEPs (*Lu et al., 2016*). However, it remains unclear whether these powdery mildew avirulence effectors bind directly to the corresponding NLR receptors or whether the receptors sense the presence of the pathogen indirectly through effector-mediated modifications of host proteins.

To reveal the molecular mechanism underlying the co-evolutionary diversification of barley MLA NLRs with host-adapted *Bgh*, we first described the natural *AVR_a* gene diversity in a local *Bgh* population. This allowed us to isolate four additional AVR_A effectors by applying a transcriptome-wide association study (TWAS) with high-confidence single nucleotide polymorphisms (SNPs) underlying disease resistance/susceptibility phenotypes in barley. We then employed co-transfection experiments to show that only matching *Mla-AVR_a* pairs activate cell death in barley leaf protoplasts. *Agrobacterium tumefaciens*-mediated co-delivery of *Mla-AVR_a* pairs in *Nicotiana benthamiana* leaves demonstrated that their co-expression is sufficient to trigger cell death in this heterologous plant species. The *AVR_a* avirulence effectors encode CSEPs that are sequence-unrelated except for allelic AVR_{A10} and AVR_{A22}. These findings are inconsistent with a co-evolutionary arms race model



describing iterative cycles of MLA receptor and AVR_A effector adaptations. Moreover, we failed to detect cell death activity for the previously reported EKA_AVR_{a10} , a member of the EKA gene family that is derived from part of a class-1 LINE retrotransposon (Ridout, 2006; Amselem et al., 2015). Using co-expression experiments with matching Mla- AVR_a pairs, including previously reported AVR_{a13} , we present evidence for direct receptor-avirulence effector interactions. Our findings imply that MLA receptors have an exceptional propensity to directly detect sequence-unrelated and likely structurally diverse pathogen effectors and that this feature might have facilitated the functional diversification of the receptor in the host population.

Results

Natural variation of AVR_a genes within a local Bgh population

In 2016 we collected 13 Bgh field isolates from leaves of barley plants within the same locality near Cologne, Germany (GPS 5°57'N, 6°51'E 5; isolates designated K2, K3, K4, S11, S15, S16, S19, S20, S21, S22, S23, S25 and S26). We purified the isolates by serial propagation of conidiospores collected from single powdery mildew colonies (Materials and methods). Then, we determined their pathotypes (virulence/avirulence profiles) together with a global collection of 17 previously described Bgh isolates (Lu et al., 2016) on a panel of near-isogenic barley lines (NILs) in the cultivar (cv.) Pallas genetic background (Kolster et al., 1986) carrying Mla1 (P01), Mla3 (P02), Mla6 (P03), Mla7 (P04B), Mla9 (P08B), Mla10 (P09), Mla12 (P10), Mla22 (P12), or Mla23 (P13) recognition specificities (Supplementary file 1). To assess infection types (ITs) independently, we additionally employed a panel of NILs in the genetic background of cv. Manchuria (Moseman, 1972) carrying introgressed Mla1 (Cl16137), Mla6 (Cl16151), Mla7 (Cl16147 and Cl16153), Mla10 (Cl16149), or Mla13 (CI16155) (Supplementary file 1). Similar to previous studies (Seeholzer et al., 2010; Lu et al., 2016), we scored five macroscopically distinguishable ITs at 9 d after conidiospore inoculation on leaves, ranging from 1 to 3 (=avirulent) or 4 to 5 (=virulent). Unambiguous assignment of virulent and avirulent interactions was possible for all isolates on Mla1, Mla6, Mla9, Mla12, Mla13, and Mla22 NILs that displayed only IT 1, 2 or 5. A subset of the Bah strains displayed IT 3 (avirulent) or 4 (virulent) on Mla3, Mla7, Mla10, and Mla23 NILs, which were in some cases difficult to distinguish from each other and thus potentially complicating the discrimination of avirulent and virulent interactions with our experimental setup (Supplementary file 1). For two of the tested Bgh isolates different avirulent/virulent interactions were recorded between Pallas and Manchuria NILs carrying Mla7 (isolates K4 and S22) and, similarly, two other isolates exhibited different avirulent/virulent interactions on Pallas and Manchuria NILs carrying Mla10 (isolates CC88 and S20; Supplementary file 1). These few discrepancies of avirulent/virulent interactions might be explained by differences in the size of introgressed chromosomal segments from the respective donor lines in Pallas and Manchuria NILs that might contain other R genes besides Mla. Therefore, we omitted the IT data of isolates K4 and S22 on MIa7 NILs and of CC88 and S20 isolates on MIa10 NILs in the TWAS (see below) to identify AVR_{a7} and AVR_{a10} candidates, respectively. Among the 13 newly described Bgh isolates collected from the same geographic site, isolates S19 and S21, as well as S23, S25, and S26 exhibited identical avirulence/virulence patterns on the NIL test panel (Supplementary file 1). Thus, 10 out of 13 isolates showed unique avirulece/virulence patterns and each isolate carries at least five different AVR_a genes (Supplementary file 1). These findings point to a high level of genetic variation among AVR_a genes within the local Bgh population.

We obtained deep fungal transcriptomes (RNA-Seq) for the ten Cologne-derived field isolates with unique avirulent/virulent patterns during pathogenesis on susceptible barley leaves at 16 and 48 hr post conidiospore inoculation, resulting in 0.8 to 6 million sequenced and mapped *Bgh* read pairs (fragments) per sample (Materials and methods). RNA-seq reads of these isolates, combined with RNA-seq reads of the previously characterized global collection of *Bgh* isolates (*Lu et al., 2016*) and of reference isolate DH14, were aligned against the recently assembled near chromosome-level DH14 genome supplemented with updated gene models (*Frantzeskakis et al., 2018*). Subsequently, a collective set of sequence polymorphisms for all isolates was identified from the combined alignment data. To assess the potential influence of population structure among the isolates prior to TWAS, we performed a population structure analysis, for which we extracted a set of high-quality diallelic and synonymous single-nucleotide polymorphisms (SNPs) from the complete set of



polymorphisms (Methods). A principal component analysis (PCA) plot and a neighbour-joining tree including all isolates confirmed the Japanese isolate RACE1 to be exceptionally divergent and assigned the three Australian isolates to a distinct clade, whereas the ten new field isolates clustered mostly together with other European, Japanese, and USA isolates (*Figure 1A*, *Figure 1—figure supplement 1*). Although all newly described isolates were collected from a single geographic location, three of these (K2, K3, and S16) appear to form a subgroup that is distinct from the other European isolates (*Figure 1A*, *Figure 1—figure supplement 1*). Interestingly, the avirulence profiles of isolates K3 and S16 on the 11 tested *Mla* barley recognition specificities belong to distinct pathotype clusters (*Figure 1B*), despite an exceptionally high similarity of K3 and S16 at the transcriptome level (*Figure 1A*).

Combined TWAS and Bgh genome analysis identified candidates for AVR_{a7} , AVR_{a9} , AVR_{a10} , and AVR_{a22}

Our TWAS analysis (*Supplementary file 2 - 4*) allowed us to identify several AVR_a candidates. Manual re-inspection of these candidates (*Lu et al., 2016*), revealed that only transcript polymorphisms of the top candidates AVR_{a7} , AVR_{a9} , AVR_{a10} , and AVR_{a22} exhibit tight linkage to the avirulence phenotypes of our Bgh strain panel on Mla7, Mla9, Mla10, and Mla22 NILs, respectively. Two genes, $BLGH_06689$ and $BLGH_06672$, encoding two identical copies of CSEP0059, were identified as topranking candidates for AVR_{a7} (p=1.78E-05, *Supplementary file 4*). This is consistent with previous reports in which two AVR_{a7} loci were inferred based on segregation analysis among the progeny of crosses between Bgh strains (*Skamnioti et al., 2008*; *Pedersen et al., 2002*). In the near chromosome-level genome assembly of isolate DH14 (*Frantzeskakis et al., 2018*) the two *CSEP0059* copies

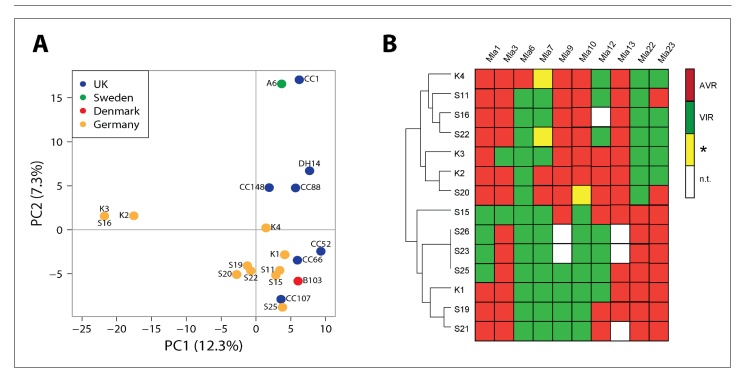


Figure 1. Population structure and avirulence profiles of *Blumeria graminis* f. sp. *hordei* (*Bgh*) isolates. (**A**) Principal Component Analysis (PCA) of the indicated European *Bgh* isolates, including ten newly collected strains from a local pathogen population in Germany, based on a set of 5170 diallelic high-quality synonymous SNPs. (**B**) Hierarchical clustering (R package 'pheatmap') of avirulence profiles from 14 *Bgh* isolates collected within the same area near Cologne, Germany (GPS 5°57'N, 6°51'E 5). Numbers correspond to infection types (ITs) 1, 2 and 3 = avirulent, red; infection types 4 and 5 = virulent, green. *denotes differences of ITs between cultivars Pallas and Manchuria, yellow. n.t.: not tested, white.

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The following figure supplement is available for figure 1:

Figure supplement 1. Population structure of a worldwide collection of 27 *Blumeria graminis* f. sp. *hordei* (*Bgh*) isolates. DOI: https://doi.org/10.7554/eLife.44471.004



are physically separated by 141 kb on scaffold 44 and, together with four other CSEPs, form a CSEP cluster in this region (Figure 2). We designated the two identical CSEP0059 copies in DH14 as candidate gene AVR_{a7}-1 (Figure 2A). One of the DH14 CSEPs in the cluster is a CSEP0059 paralog, CSEP060 (BLGH_06671), which at the protein level shares 77% aa identity with CSEP0059 (Figure 2B and C). All virulent strains on Mla7 NILs carry a SNP in CSEP0059, which results in a L51P substitution in the deduced candidate effector protein, and the corresponding variant was thus named AVR_{A7}-V1 (Figure 2A and B). Interestingly, only about half of the transcripts from the virulent isolate B103 harbor a SNP leading to the L51P substitution in the deduced protein. The other half of the B103 transcripts harbor a SNP resulting in a L28F substitution in the deduced protein (designated AVR_{A7}-V2), suggesting that in this strain the two CSEP0059 copies differ. Similarly, approximately half of the transcript reads of all three avirulent Australian isolates (Art, Aby, and Will) harbor a SNP leading to an I65T substitution in the deduced protein (designated AVR_{A7}-AUS; Figure 2A and B), suggesting that these isolates also contain an even number of non-identical CSEP0059 copies. Finally, the most divergent Bgh isolate RACE1 is clearly avirulent on Mla7 NILs (Supplementary file 1), and in its transcriptome we found two non-synonymous SNPs in all CSEP0059 transcripts corresponding to S47R and A96V aa substitutions in the deduced protein (designated AVRA7-2; Figure 2A and B). Inspection of the near chromosome-level RACE1 genome (Frantzeskakis et al., 2018) revealed two identical CSEP0059 gene copies (BGHR1_17217 and BGHR1_17236) separated by 151 kb on a single contig within a CSEP cluster that is largely syntenic with the corresponding DH14 CSEP cluster (Figure 2C; Frantzeskakis et al., 2018). RACE1 CSEP0060 (BGHR1_16533) is located on the neighbouring contig tig00005324 and is possibly syntenic to the CSEP0060 location in DH14. Additionally, RACE1 harbors a third CSEP0059 copy (BGHR1_17237) with S5N, M9T, T11A, L12W, L17F, S47R, L51P, N62S, A96V, and R105Q

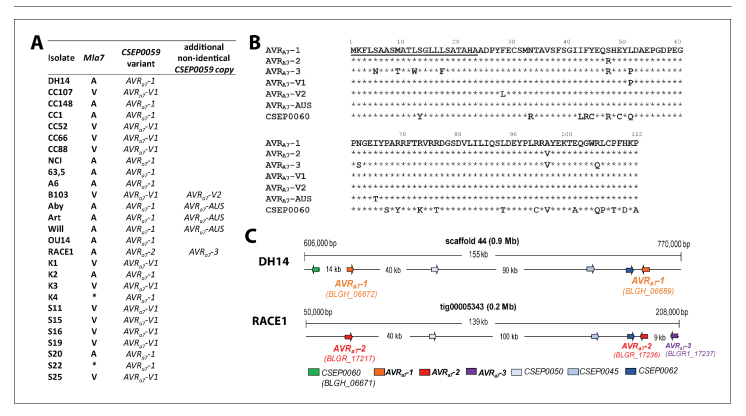


Figure 2. Identification of *CSEP0059* as an AVR_{a7} candidate by association of avirulence profiles with transcript polymorphisms and integration in the physical Bgh map. (A) AVR_{a7} transcript variants encoded by the indicated Bgh isolates with corresponding avirulence profiles. (B) Alignment of deduced AVR_{A7} amino acid sequences with all variants highlighted. (C) Visualization of the chromosomal regions harboring $CSEP0059/AVR_{a7}$ candidate variants with corresponding gene IDs in the genomes of Bgh isolates DH14 and RACE1. All CSEPs are depicted by arrows. * denotes different infection types between cultivars Pallas and Manchuria.

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substitutions in the deduced effector polypeptide (designated AVR_{A7}-3) (*Figure 2A, B and C*). Taken together, TWAS and the comparative genome analysis revealed the potential existence of multiple AVR_{A7} variants, partly encoded by paralogous genes, and two virulent AVR_{A7} variants.

TWAS identified CSEP0174 (BLGH_04994 in DH14 and BGHR1_10042 in RACE1) as the top-ranking AVR_{a9} candidate (p=1.84E-05; Supplementary file 4, Figure 3A). Analysis of the transcriptome reads showed that isolates virulent on Mla9 carry AVR_{a9} variants with either H28D (AVR_{A9}-V1) or G22E (AVR_{A9}-V2) substitutions (Figure 3A and B). Furthermore, we identified CSEP0141 as the topranking candidate for both AVR_{a10} (BGHR1_10013, p=4.50E-07) and AVR_{a22} (p=1.22E-05) (Supplementary file 4, Figure 3C). All Bgh isolates avirulent on Mla10 plants encode a CSEP0141 variant with an I77F substitution (AVR_{A10}) compared to the deduced effector protein in reference isolate DH14 (BLGH_05021; Figure 3C and D). By contrast, all isolates avirulent on Mla22 plants carry a CSEP0141 variant with 13 deduced as substitutions (V11A, F13L, D45G, D53E, Q55H, D58N, G59D, Q61P, H64Y, I77Y, V93L, W96L, and I111N; AVR_{A22} in Figure 3C and D). DH14 is the only tested isolate that is virulent on both Mla10- and Mla22-expressing plants, and we therefore named the CSEP0141 variant encoded by DH14 AVR_{a10} -V/ AVR_{a22} -V. CSEP0141 belongs to the CSEP family 64 with CSEP0266 being the only other family member that shares significant similarity with candidates AVR_{A10} and AVR_{A22} (59% and 63% identical aa sequences, respectively). Collectively, these data raised the possibility that MLA10 and MLA22 each recognize one of the two major natural variants of CSEP0141 in the *Bgh* population.

AVR_{a2} and AVR_{a10}/AVR_{a22} candidates are each single-copy genes in both DH14 and RACE1 Bgh genomes (Frantzeskakis et al., 2018). Notably, candidates AVR_{a10}-V/AVR_{a22}-V (BLGH_05021) in DH14 and AVR_{a10} (BGHR1_10013) in RACE1 physically locate approximately 300 kb away from the 3' end of AVR_{a9} (BLGH_04994 and BGHR1_10042) on the respective contiguous chromosomal DNA stretches (Figure 3E). The physical linkage of these candidate avirulence genes is consistent with a tight genetic coupling of the AVR_{a10} and AVR_{a22} loci and their linkage to the AVR_{a9} locus (Skamnioti et al., 2008). In addition, we found the transcript of candidate AVR_{a22} to map to a physical contig in the Bgh K1 genome assembly (tig19704; Hacquard et al., 2013) that is syntenic with the genomic region encompassing candidate AVR_{a10}-V/AVR_{a22}-V in DH14 and candidate AVR_{a10} in RACE1 (Figure 3E). The findings lend further support to the notion that AVR_{a10} and AVR_{a22} represent two alleles of a single Bgh effector gene (Figure 3-figure supplement 1). Candidate AVR_{a10}, which encodes a CSEP0141 variant, maps only 24 kb away from the 3' end of the only other family member CSEP0226 (BLGH_05020 and BGHR1_10014 in DH14 and RACE1 genomes, respectively) and approximately 60 kb away from the 5' end of one of the EKA class-1 LINE retrotransposon family members (Figure 3E, Amselem et al., 2015). Reminiscent of the AVRaT containing cluster of sequence-unrelated effector genes in the Bgh genome (Figure 2c), the genomic region encompassing CSEP0174 (AVR_{a9}) and CSEP0141 (AVR_{a10}/AVR_{a22}) contains sequence-unrelated CSEP0077, CSEP0080, CSEP0082, CSEP0085, CSEP0097, and CSEP0266, and therefore represents another cluster of candidate secreted effector genes in the fungal genome.

Functional analysis of AVR_a candidates in barley leaf protoplasts

We examined the capability of the identified AVR_A candidates to trigger MLA-dependent cell death upon transient co-expression with matching MLA receptors in barley leaf protoplasts. All gene constructs were expressed from the *Zea mays* ubiquitin promotor together with the firefly luciferase (LUC) reporter gene (*Lu et al., 2016*), and relative LUC activity was quantified as a proxy for protoplast viability. As epitope tag sequences can interfere with signal-noise ratios of LUC activity in this assay (*Lu et al., 2016*), we refrained from fusion of constructs with epitope sequences.

When compared to the empty vector (EV) control, co-expression of previously reported AVR_{a1} (Lu et al., 2016) with Mla1 reduced LUC activity in a Mla1-dependent manner by 71% (Figure 4A) and co-expression of AVR_{a7} -1 with Mla7 significantly reduced the reporter activity by 37% but not when co-expressed with Mla1 (Figure 4A). Interestingly, co-expression of Mla7 with either of the two AVR_{a7} variants present in RACE1, AVR_{a7} -2 and AVR_{a7} -3, resulted in a 91% and 79% reduction of LUC activity, respectively, whereas their co-expression with Mla1 did not significantly alter reporter activity (Figure 4A, Figure 4—figure supplement 1A). Taken together these data indicate that naturally occurring AVR_{a7} -variants in the Bgh population differ in their propensity to activate MLA7-mediated cell death. Neither expression of AVR_{a7} -V1 (CSEP0059) nor CSEP0060, the paralog located adjacent to AVR_{a7} -1 in the DH14 genome (Figure 3C), resulted in a significant Mla7-



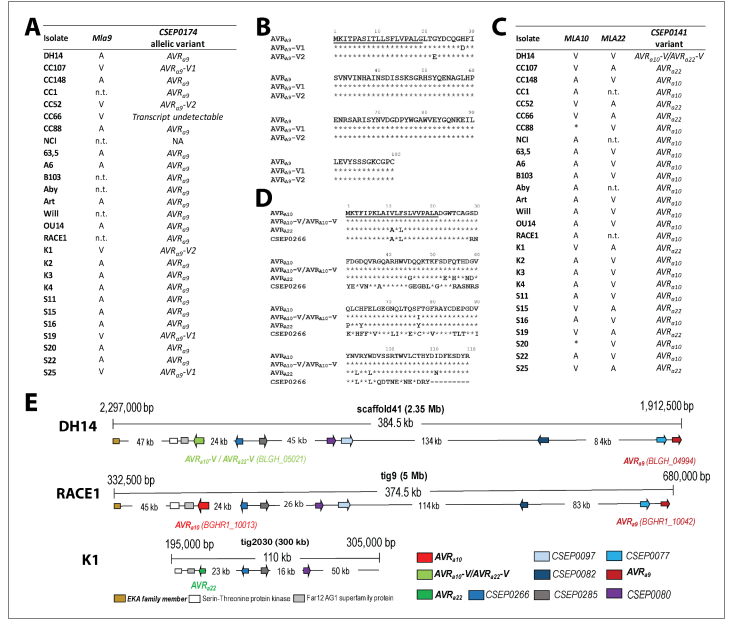


Figure 3. Identification of *CSEP0174* as an AVR_{a9} candidate and *CSEP0141* as an AVR_{a10} and AVR_{a22} candidate by association of avirulence profiles with transcript polymorphisms as well as integration in the physical Bgh map. (A) AVR_{a9} variants encoded by each Bgh isolate with corresponding avirulence profiles. (B) Alignment of deduced AVR_{A9} amino acid sequences with variants highlighted. (C) AVR_{a10}/AVR_{a22} variants encoded by each Bgh isolate with corresponding avirulence profiles. (D) Alignment of AVR_{A10} , AVR_{A22} , and AVR_{A10} - V/AVR_{A22} -V amino acid sequences. (E) Visualization of the chromosomal regions harboring $CSEP0174/AVR_{a9}$ and $CSEP0141/AVR_{a10}/AVR_{a22}$ candidates with corresponding gene IDs as well as a copy of the EKA family class-1 retrotransposon and other CSEPs in the genomes of Bgh isolates DH14, RACE1, and K1. *denotes differences of infection types between cultivars Pallas and Manchuria. n.t.: not tested.

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The following figure supplement is available for figure 3:

Figure supplement 1. Maximum likelihood phylogeny tree for the 805 predicted secreted proteins of *Bgh* DH14 lacking respective signal peptides. DOI: https://doi.org/10.7554/eLife.44471.007

dependent reduction in reporter activity (Figure 4A, Figure 4—figure supplement 1B). We detected a 30% reduction in LUC activity when AVR_{a7} -AUS was co-expressed with Mla7 but not when co-expressed with Mla1. Co-expression of AVR_{a7} -V2 with Mla7 did not result in significantly reduced LUC activity when compared to co-expression with Mla1 (Figure 4—figure supplement



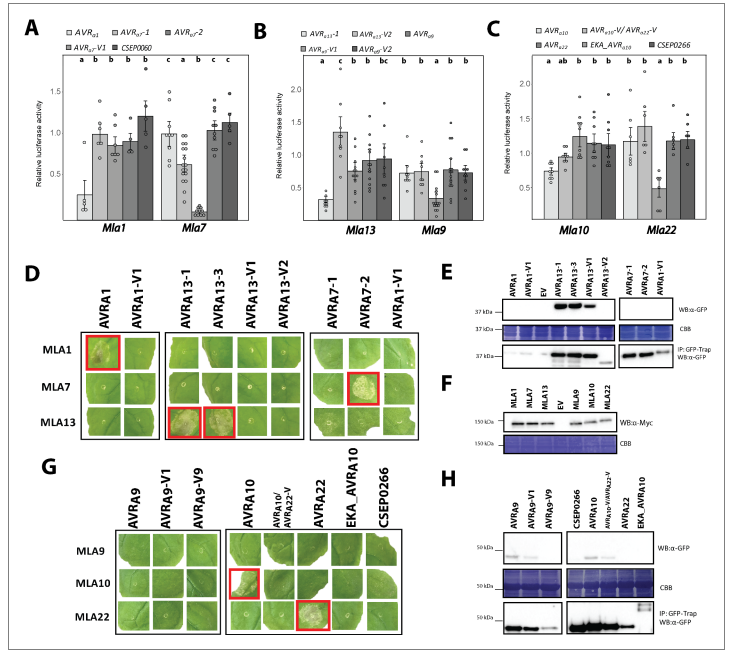


Figure 4. AVR_A candidates induce MLA-specified cell death in transient gene expression assays. (A-C) Barley protoplasts were transfected with *pUBQ: luciferase* and either an piPKb002 empty vector control (EV) or piPKb002 containing cDNAs of (A) AVR_{a1} , AVR_{a7} variants and CSEP0060, all lacking their respective signal peptides (SPs) together with either Mla1 or Mla7; (B) AVR_{a9} variants and AVR_{a13} variants, all lacking their respective SPs together with either Mla9 or Mla13; (C) AVR_{a10} , AVR_{a10} , AVR_{a22} -V, AVR_{a22} , and CSEP0266 without SPs and EKA_AVR_{a10} , together with either Mla10 or Mla22. Luciferase activity was determined ~16 hr post transfection as a proxy for cell death and normalized for each Mla construct by setting the detected luminescence for the corresponding EV transfection to 1. All values obtained in at least four independent experiments are indicated by dots, error bars = standard deviation. Differences between samples were assessed by analysis of variance (ANOVA) and subsequent Tukey post hoc tests for each Mla construct. Calculated P values were as follows: Mla7: p=1.2e-13 and Mla1: p=1.9e-03 (A); Mla13: p=3.2e-06 and Mla9: p=1.3e-04 (B); Mla10: p=4.2e-04 and Mla22: p=5.5e-04 (C). Samples marked by identical letters in the plots do not differ significantly (p<0.05) in the Tukey test for the corresponding transfected Mla. (D–H) Nicotiana benthamiana plants were transformed transiently as indicated. cDNAs lacking stop codons were fused in between the 35S promotor sequence and 4xMyc (Mla variants) or mYFP (CSEPs and AVR_a variants lacking SPs and EKA_AVR_{a10}) epitope sequences. Cell death (D, G) was determined three days post transformation and figures shown are representatives of at least three independently performed experiments with at least three transformations per experiment. Protein levels (E, F, H) of MLA-4xMyc, CSEP-mYFP, AVR_a-mYFP and EKA_AVR_a-mYFP in Nicotiana benthami



Figure 4 continued

by GFP-Trap pull-down as indicated. Extracts and immunoprecipitates were separated by gel electrophoresis and probed by anti-Myc or anti-GFP western blotting (WB) as indicated. IP: Immunoprecipitated fraction. CBB: Coomassie brilliant blue.

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The following source data and figure supplements are available for figure 4:

Source data 1. Data points indicating relative luciferase activity of Figure 4A.

DOI: https://doi.org/10.7554/eLife.44471.011

Source data 2. Data points indicating relative luciferase activity of Figure 4B.

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Source data 3. Data points indicating relative luciferase activity of Figure 4C.

DOI: https://doi.org/10.7554/eLife.44471.013

Figure supplement 1. MLA-mediated cell death in barley protoplasts.

DOI: https://doi.org/10.7554/eLife.44471.009

Figure supplement 1—source data 1. Data points indicating relative luciferase activity of Figure 4—figure supplement 1A. DOI: https://doi.org/10.7554/eLife.44471.014

Figure supplement 1—source data 2. Data points indicating relative luciferase activity of Figure 4—figure supplement 1C.

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Figure supplement 1—source data 3. Data points indicating relative luciferase activity of Figure 4—figure supplement 1D.

DOI: https://doi.org/10.7554/eLife.44471.016

Figure supplement 2. AVR_A-mYFP candidates induce MLA-4xMyc mediated cell death.

DOI: https://doi.org/10.7554/eLife.44471.010

Figure supplement 2—source data 4. Ceall death scores of samples shown in Figure 4D according to scoring system of Figure 4—figure supplement 2B.

DOI: https://doi.org/10.7554/eLife.44471.017

Figure supplement 2—source data 5. Ceall death scores of samples shown in Figure 4G according to scoring system of Figure 4—figure supplement 2B.

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1A). Barley accessions expressing the MLA7 polypeptide sequences CI1647 and CI1653 from Manchuria MLA7 NILs were found to differ slightly in two independent earlier studies (MLA7_AAQ55540, Halterman and Wise, 2004; MLA7 Seeholzer et al., 2010; Figure 4—figure supplement 1B). Therefore, we also tested the capability of MLA7_AAQ55540 to recognize AVR_{a7} and found that coexpression of $MLA7_AAQ55540$ with AVR_{a7} -2 reduced LUC activity by only 68% (as compared to 91% when co-expressed with Mla7). Luciferase activity in protoplasts co-expressing $MLA7_AAQ55540$ and AVR_{a7} -1 (30% LUC reduction compared to EV) did not differ significantly from protoplasts co-expressing $MLA7_AAQ55540$ and AVR_{a7} -V1 (15% LUC reduction compared to EV; Figure 4—figure supplement 1C).

Co-expression of Mla9 with the AVR_{a9} candidate, but not its variants AVR_{a9} -V1 and AVR_{a9} -V2 or AVR_{a13} -V1 and AVR_{a13} -V2 (Lu et al., 2016), resulted in a significant 67% relative reduction of LUC activity. This cell death activity was specific to Mla9 as co-expression of AVR_{a9} with Mla13 did not significantly reduce LUC activity (Figure 4B). Co-expression of the previously reported AVR_{a13} (Lu et al., 2016) with Mla13 reduced LUC activity in a Mla13-dependent manner by 69% (Figure 4B).

Similar to the moderate reduction of LUC activity when AVR_{a7} -1 was co-expressed with Mla7 (**Figure 4A**), co-expression of the AVR_{a10} candidate with Mla10 reduced the reporter activity by only 25% (**Figure 4C**). LUC reduction was not observed when AVR_{a10} - V/AVR_{a22} -V, AVR_{a22} , CSEP0266, or EKA_AVR_{a10} were co-expressed with Mla10. Co-expression of EKA_AVR_{a10} , AVR_{a10} , and Mla10 had no impact on LUC reduction when compared to co-expression of AVR_{a10} and Mla10 alone (**Figure 4—figure supplement 2D**). Co-expression of Mla22 with AVR_{a22} , but not AVR_{a10} , AVR_{a10} - V/AVR_{a22} -V, CSEP0266, or EKA_AVR_{a10} , reduced relative LUC activity by 53% when compared to co-expression of Mla22 with an EV control (**Figure 4C**). These findings provide functional evidence that CSEP0141 alleles define AVR_{a10} and AVR_{a22} avirulence effectors and are specifically recognized by allelic MLA10 and MLA22 receptors, respectively.



Co-expression of matching Mla and AVR_a pairs is necessary and sufficient to trigger cell death in N. benthamiana

Next we tested whether *Agrobacterium tumefaciens*-mediated delivery and co-expression of matching *Mla* and *AVR_a* pairs can trigger cell death in heterologous *N. benthamiana* leaves. In addition to newly isolated AVR_{A7}, AVR_{A9} and AVR_{A10}/AVR_{A22}, we included the previously reported AVR_{A1} and AVR_{A13} variants as additional specificity controls in these experiments (*Figure 4D–4H*) (*Lu et al.*, *2016*). For protein expression and stability analysis in this heterologous system, the constructs were designed to express C-terminally 4xMyc-tagged MLA receptors and C-terminally mYFP-tagged AVR_A variants without signal peptide sequences.

Delivery of the AVR_{a1}-mYFP construct but not the construct for its virulent variant, AVR_{a1}-V1-mYFP, conferred cell death in N. benthamiana when co-expressed with Mla1-4xMyc, but not when co-expressed with Mla7-4xMyc or Mla13-4xMyc (Figure 4D, Figure 4—figure supplement 2). We also observed statistically significant cell death intensity when AVR_{a13}-1-mYFP and AVR_{a13}-3-mYFP, but not the virulent variants AVR_{a13}-V1-mYFP or AVR_{a13}-V2-mYFP when co-expressed with Mla13-4xMyc (Figure 4D; Figure 4—figure supplement 2). Cell death was not seen when the same AVR_a effector constructs were co-expressed with Mla1-4xMyc or Mla7-4xMyc (Figure 4D; Figure 4—figure supplement 2), indicating retained recognition specificity of MLA1 and MLA13 receptors in this heterologous plant species, respectively.

 AVR_{a7} -1 mediates moderately reduced LUC reporter activity in barley protoplasts when co-expressed with Mla7, whereas AVR_{a7} -2 expression leads to a strong reduction of reporter activity in the same experiment (**Figure 4A**). Correspondingly, in N. benthamiana we observed MLA7-dependent cell death only when expressing AVR_{a7} -2-mYFP together with Mla7-4xMyc, but not when Mla7-4xMyc was co-expressed with AVR_{a7} -1-mYFP or AVR_{a7} -V1-mYFP variants (**Figure 4D**, **Figure 4—figure supplement 2**). The lack of AVR_{a7} -1 - MLA7 cell death in N. benthamiana is not due to AVR_{a7} -1-mYFP protein stability (**Figure 4E**) but may be due to other unknown aspects of the heterologous system.

Major differences in protein steady-state levels were found between individual AVR_A effectors, whereas protein levels of all MLA receptors were comparable in α -Myc western blots (*Figure 4E and F*). mYFP-tagged AVR_{A1} variants were barely detectable even after enrichment by GFP-Trap (*Figure 4E*), whilst AVR_{A13}-1, AVR_{A13}-3, and AVR_{A13}-V2 were detectable in *N. benthamiana* extracts without GFP-Trap pull-down (*Figure 4E*). AVR_{A13}-V1-mYFP protein was barely detectable even after GFP-Trap enrichment (*Figure 4E*) suggesting that loss of MLA13-mediated cell death activity for AVR_{A13}-V1 may be due to protein instability. Notably, we found a Western blot signal corresponding to the expected size of ~40 kDa for the AVR_{A13}-V2-mYFP fusion protein (*Figure 4E*). This differs from the expression of *FLAG-AVR*_{a13}-V2 in barley protoplasts, where most of the FLAG-AVR_{A13}-V2 fusion protein was visible as a cleaved protein product (*Lu et al., 2016*). mYFP-tagged AVR_{A7} variants were only detectable in *N. benthamiana* leaf extracts after GFP-Trap pull-downs (*Figure 4E*).

 AVR_{a9} elicited a 67% reduction in LUC activity when co-expressed with Mla9 in barley protoplasts (*Figure 4B*). Surprisingly however, in N. benthamiana, neither AVR_{A9} -mYFP nor its virulent variants AVR_{A9} -V1-mYFP or AVR_{A9} -V2-mYFP triggered MLA9-dependent cell death (*Figure 4G*; *Figure 4*—figure supplement 2) although mYFP-tagged AVR_{A9} was detectable in N. benthamiana protein extracts (*Figure 4H*). We postulate either that a third barley protein other than MLA9 and AVR_{A9} is needed for MLA9-mediated cell death activation in N. benthamiana or that the chosen combination of epitope tags prevents effective AVR_{A9} recognition in this plant species.

Although AVR_{a10} reduced LUC reporter activity in barley protoplasts only moderately when co-expressed with Mla10 (Figure 4C), we detected a clearly visible cell death response when AVR_{a10}-mYFP was co-expressed with Mla10-4xMyc in N. benthamiana leaves in multiple independent experiments (Figure 4G; Figure 4—figure supplement 2). The MLA10-triggered cell death specifically required the presence of AVR_{A10}-mYFP because leaf cells remained alive upon co-expression with AVR_{A10}-V/AVR_{A22}-V-mYFP or AVR_{A22}-mYFP (Figure 4G; Figure 4—figure supplement 2). Consistent with data obtained with barley leaf protoplasts, cell death in heterologous N. benthamiana was induced upon co-expression of Mla22-4xMyc with AVR_{a22}-mYFP, but not AVR_{a10}-mYFP or AVR_{a10}-V/AVR_{a22}-V-mYFP (Figure 4G; Figure 4—figure supplement 2). Furthermore, transient expression of CSEP0266-mYFP or EKA_AVR_{a10}-mYFP together with either Mla10-4xMyc or Mla22-4xMyc failed to trigger cell death in N. benthamiana (Figure 4G; Figure 4—figure supplement 2), corroborating



our conclusion that allelic AVR_{A10} and AVR_{A22} avirulence effectors are specifically recognized by allelic MLA10 and MLA22 receptors, respectively. Only AVR_{A10}-mYFP and AVR_{A10}-V/AVR_{A22}-V-mYFP were detectable in *N. benthamiana* protein extracts and these two fusion proteins as well as CSEP0266-mYFP and AVR_{A22}-mYFP ran at the expected size of ~40 kDa after enrichment by GFP-Trap pull-downs (*Figure 4H*). In these experiments the AVR_{A22}-mYFP protein appears less stable than AVR_{A10}-mYFP (*Figure 4H*). EKA_AVR_{A10}-mYFP is expected to migrate at ~70 kDa. We detected a western blot signal at this expected size and one faster-migrating variant (*Figure 4H*). In conclusion, we failed to detect cell death activity in response to the previously reported recognition of EKA_AVR_{a10} by MLA10 (*Ridout, 2006*). This contrasts with the functional validation of the AVR_{a10} candidate identified here by TWAS.

Candidate AVR_A proteins interact with matching MLA receptors in plant extracts and in yeast

AVR_{A1} is recognised by MLA1 in barley, A. thaliana and, as shown here, also in N. benthamiana (Figure 4D; Lu et al., 2016). The retention of MLA1-dependent recognition of AVR_{A1} in three divergent plant families (Triticeae, Brassicaceae, Solanaceae) suggests direct interactions of matching MLA and AVRA pairs or an indirect recognition mechanism involving highly evolutionarily conserved AVR_A host target(s). The wheat Mla ortholog Sr50 interacts with its cognate effector AvrSr50 of Puccinia graminis f. sp. tritici in yeast (Chen et al., 2017). Despite the lack of sequence conservation between most of the identified AVRA effectors and AvrSr50, we tested whether barley MLA directly interacts with cognate Bgh effectors. So far it has been impossible to purify large quantities of recombinant full-length MLA receptors for in vitro AVRA-MLA association studies, possibly because of MLA-triggered cell death and receptor oligomerisation (Maekawa et al., 2011b). We thus focused on quantitatively measuring putative AVRA-MLA associations in plant extracts using the highly sensitive split-luciferase (split-LUC) complementation assay (Paulmurugan et al., 2002; Luker et al., 2004). Whereas barley protoplasts can undergo cell death upon expression of matching Mla and AVRa pairs at ~16 hr post transfection (Figure 4A-4C), expression in N. benthamiana leaves permitted MLA and AVRA interaction analysis at two days post A. tumefaciens leaf infiltration and prior to the appearance of macroscopically visible MLA-mediated cell death. To examine MLA and AVRA associations by luciferase activity of protein extracts from A. tumefaciens-infiltrated leaf area, we generated gene constructs in which MLA and AVR_A were fused at the C-terminus to cLUC and nLUC, respectively (Mla-cLUC and AVRa-nLUC), and used these for A. tumefaciens-mediated transient gene expression experiments in N. benthamiana leaves (Materials and methods). We focused on AVR_{a13}, AVR_{a7}, and AVR_{a10}/AVR_{a22} variants and their cognate Mla receptors for split-LUC assays (Figure 5A-5C), because AVR_{A1} protein levels were barely detectable (Figure 4E), and co-expression of Mla9 and AVRa9 failed to trigger a cell death response in N. benthamiana leaves (Figure 4G).

We detected high LUC activities (>10,000 units) in extracts of leaves expressing Mla13-cLUC together with $AVR_{a13}\text{-}1\text{-}nLUC$ and $AVR_{a13}\text{-}3\text{-}nLUC$ but not when Mla13-cLUC was exchanged to Mla1-cLUC (Figure 5A), although the two MLA-cLUC proteins were similarly stable (Figure 5D). This was not the case for samples expressing $AVR_{a13}\text{-}V1\text{-}nLUC$, possibly due to lack of detectable $AVR_{a13}\text{-}V1\text{-}nLUC$ protein (Figure 5E). Unexpectedly, the highest LUC activity was seen when Mla13-cLUC was expressed together with $AVR_{a13}\text{-}V2\text{-}nLUC$ (>100,000 units; Figure 5A). The high LUC activities were dependent on the MLA13 receptor because only low LUC activities (<300 units) were observed when the same AVR_{a13} variants were co-expressed with Mla1-cLUC (Figure 5A).

We detected LUC activity when co-expressing Mla7-cLUC with AVR_{a7} -2-nLUC, which was at least 7-fold higher when the latter construct was replaced by AVR_{a7} -1-nLUC or AVR_{a7} -V1-nLUC (1000 units; **Figure 5B**), despite comparable protein levels of all AVR_{A7} -nLUC variants (**Figure 5E**). The LUC activities were much lower compared to the LUC complementation of Mla13-cLUC and AVR_{a13} -1-nLUC (**Figure 5A**). The higher LUC reporter activity was not detected when co-expressing Mla13-cLUC with AVR_{a7} -2-nLUC. This proxy for in planta receptor-avirulence effector interaction is in agreement with our observation that AVR_{a7} -2-mYFP but not AVR_{a7} -1-mYFP or AVR_{a7} -V1-mYFP is capable of inducing MLA7-dependent cell death in N. benthamiana (**Figure 4D**).

Co-expression of AVR_{a10} -nLUC with Mla10-cLUC resulted in the detection of a 3.5-fold higher LUC activity when compared to expression of its virulent variants AVR_{a10} - V/AVR_{a22} -V-nLUC and AVR_{a22} -nLUC (Figure 5C). LUC activity upon expression of AVR_{a22} -nLUC together with Mla22-cLUC



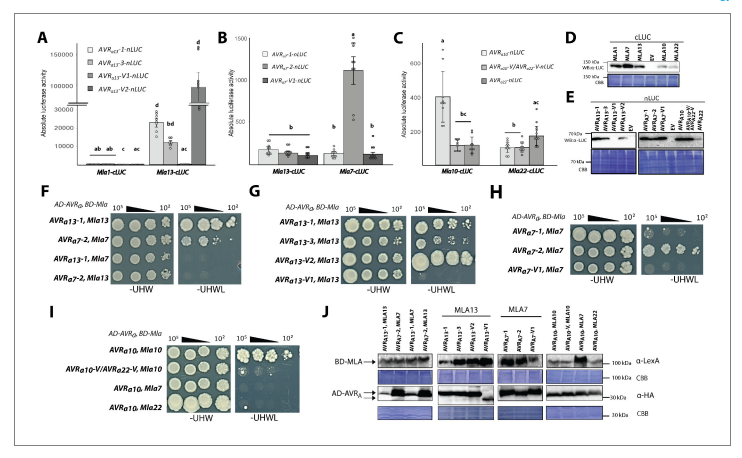


Figure 5. Association of candidate AVRA proteins with MLA in plant extracts (A-E) and in yeast (F-J). (A-C) Nicotiana benthamiana plants were transformed transiently with constructs encoding (A) Mla1-cLUC or Mla13-cLUC together with cDNAs of AVR_{a13}-1, AVR_{a13}-3, AVR_{a13}-V1, AVR_{a13}-V2 lacking signal peptides (SPs) and fused C-terminally in frame with nLUC, (B) Mla1-cLUC, or Mla7-cLUC together with cDNAs of AVRa7-1, AVRa7-2, and AVR_{a7}-V1 lacking SPs and fused C-terminally in frame with nLUC, (C) Mla10-cLUC or Mla22-cLUC together with cDNAs of AVR_{a10}, AVR_{a10}-V/AVR_{a22}-V, and AVR_{a22} without SPs fused C-terminally in frame with nLUC, all under the control of the 35S promotor. Luciferase activity was determined two days post transfection. All values obtained in at least six experiments are indicated by dots, error bars = standard deviation. For each graph, differences between samples were assessed using non-paramatric analysis of variance (Kruskal-Wallis) and subsequent Dunn's post hoc tests. Calculated P values were as follows: (A) p=6.8e-10, (B) p=1.2e-04, (C) p=8.0e-07. Samples marked by identical letters in the plot did not differ significantly (p<0.05) in Dunn's test. (D-E) Protein levels of MLA-cLUC (D) and AVR_A-nLUC (E) variants in Nicotiana benthamiana corresponding to constructs of 5A to 5C. Leaf tissue was harvested two days post infiltration. Total protein was extracted, separated by gel electrophoresis and probed by anti-LUC western blotting (WB). (E-I) Yeast cells were co-transformed with Mla alleles fused N-terminally to the LexA binding domain (BD) and AVRa constructs lacking SPs fused N-terminally to the B42 activation domain (AD) and 1xHA tag as indicated. Growth of transformants was determined on selective growth media containing raffinose and galactose as carbon sources but lacking uracil, histidine and tryptophan (-UHW), and interaction of proteins was determined by leucine reporter activity reflected by growth of yeast on selective media containing raffinose and galactose as carbon sources but lacking uracil, histidine, tryptophan and leucine (-UHWL). Figures shown are representatives of at least three independent experiments with yeast clones obtained from three independent yeast transformation experiments and pictures were taken 12 days after drop out. (J) Protein levels of BD-MLA and AD-AVRA variants corresponding to yeast of Figure 5D-5G. Yeast transformants were grown in glucose containing selective media lacking uracil, tryptophan, and histidine to OD₆₀₀ = 1. Cells were harvested, total protein extracted, separated by gel electrophoresis, and western blots (WB) were probed with anti-LexA or anti-HA antibodies as indicated.

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The following source data is available for figure 5:

Source data 1. Data points indicating absolute luciferase activity of Figure 5A.

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Source data 2. Data points indicating absolute luciferase activity of Figure 5B.

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Source data 3. Data points indicating absolute luciferase activity of Figure 5C.

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was only slightly higher (1.6-fold) when compared to its virulent variants AVR_{a10} -nLUC and AVR_{a10} - V/AVR_{a22} -V-nLUC in the same experiment (**Figure 5C**). This marginal increase in LUC activity may partly reflect differences in protein stability, as the AVR_{A22} -nLUC protein is barely detectable when compared to AVR_{A10} -nLUC (**Figure 5E**) and because both, MLA10-nLUC and MLA22-nLUC are seemingly less stable when compared to MLA7-nLUC and MLA13-nLUC.

Although the LUC complementation assay is suggestive of direct receptor - avirulence effector associations, we cannot fully exclude the involvement of other plant proteins in this association. We thus tested MLA and AVRA interactions in a yeast two-hybrid assay using leucine reporter gene activity (Figure 5F-5I). Similar to results obtained with the split-LUC assay, yeast growth on leucine-deficient medium was observed when we co-expressed AD (B42 activation domain)-AVR_{a13}-1 with BD (LexA binding domain)-Mla13 and AD-AVR_{a7}-2 with BD-Mla7, but not when BD-Mla13 and BD-Mla7 were swapped in these two interaction experiments, indicating the interactions are specific for matching MLA receptor and avirulence effector pairs (Figure 5F). Interactions in yeast were detectable when BD-Mla13 was co-expressed with either AVR_{a13} -1 or AVR_{a13} -3, but undetectable upon coexpression with AVR_{a13}-V1, suggesting specific MLA13 interactions with the avirulent AVR_{A13} variants (Figure 5G). However, clear interactions in yeast were also detectable upon co-expression of BD-Mla13 with AD-AVR_{a13}-V2, even at a cell plating density of 10² (Figure 5E), which is reminiscent of the split-LUC result with the corresponding construct pair (Figure 5A). In contrast to AVR_{A13}-V1mYFP (Figure 4E) and AVR_{A13}-V1-nLUC (Figure 5E) protein in N. benthamiana leaves, AD-AVR_{A13}-V1 protein level in yeast was comparable to other AD-AVR_{A13} variants (Figure 5J). Robust interaction in yeast was found upon co-expression of BD-Mla7 with AD-AVR_{a7}-2, but not when the former was co-expressed with AD-AVR_{a7}-V1 (Figure 5H). Only sporadic yeast colony growth was found when BD-Mla7 was co-expressed with AD-AVR_{a7}-1 (Figure 5H). This again mirrors the findings with the corresponding gene pairs in the LUC complementation assay (Figure 5B). We conclude that specific interactions can be detected for the matching MLA7 and AVR_{A7}-2 pair in yeast and specific associations for the corresponding protein pair in plant extracts. In sum, yeast two-hybrid and split-LUC experiments suggest direct detection of the sequence-unrelated avirulence effectors AVR_{A7} and AVR_{A13} by matching MLA7 and MLA13 receptors, but the strong association of the virulent effector AVR_{A13}-V2 with MLA13 represents one case in which receptor-effector association is uncoupled from receptor activation, that is from cell death induction (see Discussion).

Consistent with a direct interaction of matching MLA and AVR_A pairs, co-expression of *BD-Mla10* with $AD-AVR_{a10}$ in yeast resulted in leucine reporter gene activation and this was undetectable when *Mla10* was replaced by either *Mla2* or *Mla22*, and minor when $AD-AVR_{a10}$ was replaced by its virulent variant $AD-AVR_{a10}-V/AVR_{a22}-V$ (*Figure 5I*). The latter virulent effector differs only by a single amino acid from the avirulence effector AD-AVR_{A10} (*Figure 3D*).

Evolutionary history of *Bgh AVR*_a genes and population-level *AVR*_{a10}/*AVR*_{a22} sequence variation in *B. graminis formae speciales*

We used a high-quality genome assembly of the wheat powdery mildew Bgt reference isolate 96224 to investigate the evolutionary history of Bgh AVR_a genes and to potentially identify distinctive selection pressures on these effector genes exerted by wheat and barley hosts. AVR_{a7} (sequence identical $BLGH_06672$ and $BLGH_06689$; **Figure 2**) belongs to a larger CSEP gene family in the Bgh DH14 genome and, together with three closely related members, defines one sublineage (**Figure 6—figure supplement 1A**). Although we identified four genes sharing between 40% and 47% as sequence similarity with AVR_{a7} (**Figure 6—figure supplement 1A**), the flanking genes of AVR_{a7} differ from those adjacent to the four Bgt homologs and therefore do not permit conclusions on potential orthologous relationships after the split of the two *formae speciales*.

We found three *Bgt* genes sharing between 52% and 57% identical polypeptide sequences with *Bgh AVR_{a9}* (*BLGH_04994*) (*Figure 6—figure supplement 1C and D*). These genes are closely located to each other in a region of the wheat powdery mildew genome that is largely collinear between *Bgt* and *Bgh* genomes (*Figure 6—figure supplement 1E and F*). We conclude that either local gene duplications of an ancestral ortholog of *BLGH_04994* gave rise to the extant *Bgt* gene organization, or that these duplications were already present in the last common ancestor of *Bgt* and *Bgh* and that paralogs were lost in the *Bgh* genome.

Applying a phylogenetic approach to the wheat powdery mildew genome, we identified BgtE-5921 as the ortholog of Bgh CSEP0141 (with haplotypes AVR_{a10} and AVR_{a22}), which shares 68% and



65% identical deduced polypeptide sequences with AVR_{A10} and AVR_{A22}, respectively (Figure 6—figure supplement 1G; Table 1; Figure 6A). In addition, we found a single CSEP0141 candidate ortholog in each of the genomes of a world-wide collection of other B. graminis formae speciales (Table 1, Supplementary file 5), for which short-read genome sequences are available (Spanu et al., 2010; Wicker et al., 2013; Hacquard et al., 2013; Menardo et al., 2016; Praz et al., 2017; Müller et al., 2019). With these short-read genome sequences from multiple isolates of each of these B. graminis f. sp. (Supplementary file 5) we were able to assess genome-wide nucleotide diversity (π) and population-level sequence diversification of the CSEP0141 orthologs and compared them with the diversification pattern of CSEP0141 in the Bgh population. Based on 1141 neutral markers, we calculated a genome-wide per-site nucleotide diversity (π) of 0.022 for the f. sp. hordei, 0.013 for the f. sp. secalis, 0.050 for the f. sp. tritici2, 0.062 for the f. sp. tritici, and 0.040 for the f. sp. triticale isolates (Supplementary file 5). Unlike the two major CSEP0141 haplotypes in Bgh (AVR_{a10} and AVR_{a22}), a single dominant BgtE-5921 haplotype was found in the Bgt population (29 out of 40 isolates; Table 1, Figure 6A). The remaining 11 Bqt isolates represent six further BqtE-5921 haplotypes encoding effector variants with at most four aa polymorphisms in the deduced proteins compared to the dominant BgtE-5921 variant (Table 1). Three of the latter haplotypes are exclusively present in Bgt isolates that were collected from tetraploid wheat and represent a distinctive Bgt sublineage, designated B. graminis f. sp. tritici2 (Menardo et al., 2016). A single haplotype of the CSEP0141 ortholog was detected among 22 B. graminis f. sp. triticale isolates and a single haplotype was also found in five B. graminis f. sp. secalis isolates (Table 1), indicating either limited or no natural variation of this effector in populations of wheat, triticale, and rye powdery mildews (up to four as substitutions in wheat powdery mildews). This underlines the exceptional level of polymorphism between the two major CSEP0141 haplotypes, AVR_{a10} and AVR_{a22} , in the Bgh population with 13 deduced as changes. Bgh CSEP0141 defines one of 190 core effector genes that are conserved across B. graminis ff. spp. (Frantzeskakis et al., 2018) and exhibits the highest frequency of non-synonymous SNPs in the Bgh population (4.2 non-synonymous SNPs/100 bp coding sequence; Figure 6C). Virulence functions of core effectors are likely important for B. graminis pathogen fitness. Thus, we hypothesize that in the Bgh population the two dominant CSEP0141 haplotypes, AVR_{a10} and AVR_{a22} , have emerged from the action of two opposing selective pressures: sequence conservation that maintains pathogen fitness through retention of effector virulence activity and sequence diversification to escape recognition by MLA10 and MLA22 receptors, respectively.

Discussion

Previous pathotyping studies of *Bgh* field isolates with barley varieties carrying different powdery mildew *R* genes suggested that the Central European pathogen population can be considered as a single epidemiological unit (*Limpert, 1987*). We have shown here that among 13 *Bgh* isolates from

Table 1. AVR_{a10}/AVR_{a22} ortholog BgtE-5921 variants in different Blumeria graminis formae speciales.

	Number of	Number of variants in Blumeria graminis f. sp.				
AVR _{a10} /AVR _{a22} ortholog BgtE-5921	tritici	dicocci/tritici2	triticale	secalis	Total	
Нар_96224	29		22		51	
Hap_96224 R111G	2				2	
Hap_96224 V5I, R111G	2				2	
Hap_96224 R111G, A115G	1				1	
Hap_96224 F77L, R111G		4			4	
Hap_96224 F77L, R111G, Y117H		1			1	
Hap_96224 V5I, I8L, F77L, R111G		1			1	
Hap_96224 V17I, S47M, D48N, R50K, G58D, G59S, R101C, R111G, A115V				5	5	
total number of isolates	34	6	22	5	67	

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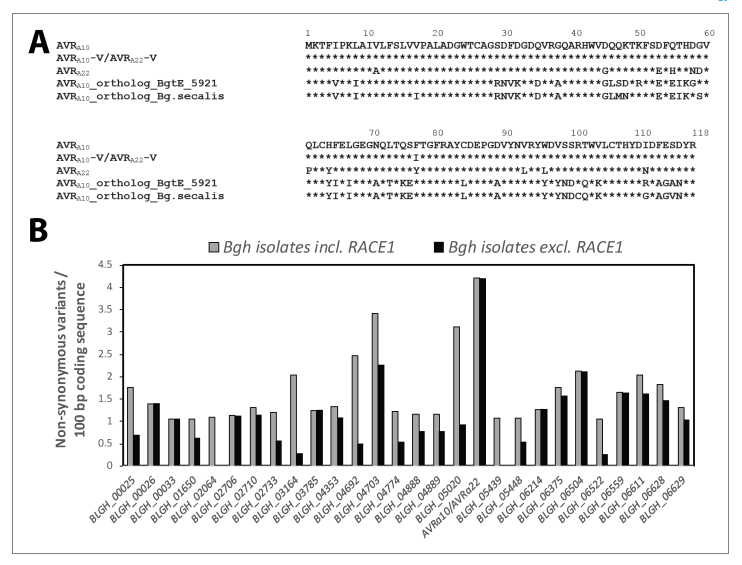


Figure 6. Conservation of AVR_{a10}/AVR_{a22} orthologs between *Blumeria graminis formae speciales*. (A) Alignment of protein sequences (AVR_{A10}/AVR_{A22}) encoded by *Bgh CSEP014*1 and orthologs detected in *B. graminis* f. sp. *tritici* and *B. graminis* f. sp. *secalis*. (B) Number of non-synonymous sequence variants was determined for 190 core effectors (*Frantzeskakis et al., 2018*) among all *Bgh* isolates described in this study and is displayed for all core effectors with ≥1 non-synonymous variants/100 bp coding sequence. Grey bars, including all *Bgh* isolates; black bars, all *Bgh* isolates excluding RACE1. DOI: https://doi.org/10.7554/eLife.44471.023

The following figure supplement is available for figure 6:

Figure supplement 1. Phylogenetic and genomic analysis of AVR_{a7}, AVR_{a9}, AVR_{a10}/AVR_{a22} homologs in *Blumeria graminis* f. sp. tritici. DOI: https://doi.org/10.7554/eLife.44471.024

a local population in Germany ten virulence combinations can be distinguished in interactions with a panel of Mla NILs, suggesting potential adaptation of the pathogen population to multiple Mla resistance specificities. Our findings are congruent with a recent study describing a high complexity of Central European Bgh virulence pathotypes on a panel of 50 differential barley varieties carrying Mla or other powdery mildew R genes (Dreiseitl, 2014). The same study also provided evidence for a complete separation of the Central European and Australian Bgh populations with non-overlapping pathotypes and estimated an almost three-fold higher virulence complexity for the European population. This is believed to be due to the cultivation of barley varieties carrying distinct powdery mildew R gene combinations on the two continents, which leads to differential intensities of directed selection on Bgh populations. Analysis of our isolate collection comprising, among others, 13 newly characterized Bgh strains from a local population in Germany and three representative Australian



isolates, is supportive of a significantly greater virulence complexity of the Central European over the Australian isolates even when considering virulence patterns only on *Mla* NILs (*Supplementary file 1*). The complexity of avirulence and virulence alleles in the Central European population suggests that the fungus adapts by balancing selection of *AVR*_a genes among strains rather than selective sweeps to maintain pathogenicity.

Specific activation of cell death upon transient gene expression of matching AVR_a and Mla gene pairs in barley protoplasts and heterologous N. benthamiana provided evidence for the function of candidate AVR_{a7}, AVR_{a9}, AVR_{a10}, and AVR_{a22} as avirulence effectors, all of which encode Bgh CSEPs with a predicted signal peptide. This is different from the previously reported EKA_AVR_{A10}, which lacks a canonical signal peptide, belongs to the large EKA gene family that is derived from part of a class-1 LINE retrotransposon (Ridout, 2006; Amselem et al., 2015), and was used as an elicitor of induced MLA10 nuclear interaction with a WRKY transcription factor (Shen et al., 2007). However, we were unable to detect MLA10-mediated cell death activity for EKA_AVR_{A10}. Our findings demonstrate that virulent Bgh isolates escape recognition of corresponding MLA receptors predominantly by non-synonymous SNPs but also loss of expression of the corresponding genes (Figure 3A; Lu et al., 2016). Upon in silico removal of the signal peptide, phylogenetic analysis for the 805 predicted secreted proteins of Bgh (Frantzeskakis et al., 2018) and comparative sequence analysis of AVR_{a7}, AVR_{a9}, AVR_{a10}, AVR_{a22}, and previously reported AVR_{a1} and AVR_{a13}, also representing CSEPs, failed to detect evolutionary conservation (Figure 3-figure supplement 1) or significant sequence similarity (at most 8% sequence identity) between any pair of these polypeptides except for allelic AVR_{a10} and AVR_{a22} (Figure 3C and E). By contrast, MLA7, MLA9, MLA10 and MLA13 receptors are >96% identical in aa sequence to each other, whereas MLA1 and MLA22 are more diverged and share 91% identical polypeptide sequences with this receptor group (Seeholzer et al., 2010). This raises questions regarding the evolutionary history of MLA10- AVR_{A10} and MLA22-AVR_{A22} receptoreffector pairs. AVR_{a10} and AVR_{a22} are notable for several reasons: (i) these allelic effectors define two dominant haplotypes of CSEP0141 in Bgh, (ii) CSEP0141 belongs to a core of 190 effectors that are conserved among different B. graminis f. sp. and, therefore, likely contributes to pathogen fitness, (iii) CSEP0141 represents the core effector with the highest frequency of non-synonymous SNPs among the tested global collection of Bgh isolates, and (iv) a single dominant haplotype of its Bgt ortholog, designated BgtE-5921, was found in the wheat powdery mildew population. Collectively, this suggests a model in which the two dominant CSEP0141 haplotypes, AVR_{a10} and AVR_{a22}, evolved in Bgh by the action of two opposing selective forces: functional conservation that maintains pathogen fitness through retention of effector virulence activity and sequence diversification to escape recognition by MLA10 and MLA22 receptors, respectively. AVR_{a10} and AVR_{a22} likely represent a balanced polymorphism in the extant pathogen population because Bgh isolates containing one or the other haplotype do not form discrete subgroups (Figure 1-figure supplement 1A). However, the likely source of many, if not all, Mla recognition specificities in domesticated barley is wild barley, H. spontaneum (Jørgensen and Wolfe, 1994). Thus, it is possible that Mla10, Mla22 and their matching avirulence effector genes have not diversified during a co-evolutionary arms race (Ravensdale et al., 2011) but have rather evolved independently in separate host and pathogen populations. In such a scenario, the balanced AVR_{A10} and AVR_{A22} polymorphism in the extant pathogen population is the consequence of concurrent cultivation of domesticated barley varieties with these Mla resistance specificities. Besides an apparently balanced AVR_{a10} and AVR_{a22} polymorphism at CSEP0141 in the Bgh field population one would expect sporadic strains that are virulent on both Mla10- and Mla22-harboring host varieties, which is the case for Bgh isolate DH14 carrying a SNP that introduces a single aa substitution in AVR_{a10} (Figure 3). In such strains, a fitness penalty for the pathogen might be mitigated by residual virulence activity of the CSEP0141 variant.

Our split-LUC and yeast two-hybrid experiments provided evidence for direct and specific interactions between multiple matching MLA/AVR_A pairs (*Figure 5*). More than 50 years after the original discovery of multi-allelic race-specific disease resistance at *Mla* (*Moseman and Schaller, 1960*), these findings now imply that the co-evolutionary functional diversification of these immune receptors is at least in part mechanistically underpinned, and was perhaps driven by direct interactions with sequence-unrelated *Bgh* avirulence effectors. Direct receptor-avirulence effector interactions have been described for flax L and rice Pik multi-allelic disease resistance genes, which encode NLR-type receptors, (*Kanzaki et al., 2012; Dodds et al., 2006*). In flax, a subset of allelic L receptors (L5, L6, and L7) interact with a specific subset of highly sequence-related flax rust AvrL567 proteins, and



in rice, allelic Pik receptors interact with specific variants of highly sequence-related AVR-Pik proteins. Rice Pik immune receptors contain an integrated heavy metal-associated RATX1/HMA domain, which binds directly to AVR-Pik and Pik functional diversification is driven by polymorphisms in this integrated domain (Kanzaki et al., 2012; Magbool et al., 2015; De la Concepcion et al., 2018). Barley MLA and flax L proteins lack detectable integrated domains, and diversifying selection in allelic MLA receptors is largely confined to predicted solvent-exposed residues of leucine-rich repeats 7 to 15 (Seeholzer et al., 2010). Thus, to the best of our knowledge, a co-evolutionary functional diversification of multi-allelic NLR-type receptors in plants with directly recognized sequence-unrelated avirulence effectors, as described here for matching MLA, is without precedence. Effectors AVR1-CO39 and AVR-Pia of the ascomycete Magnaporthe oryzae are sequence-unrelated but have very similar 6 β -sandwich structures that are stabilized in both cases by a disulfide bridge and are both recognized by the rice NLR pair RGA4-RGA5 through the integrated RATX1/HMA domain located at the C-terminus of RGA5 (de Guillen et al., 2015; Cesari et al., 2013). Structural similarity searches then showed that AVR1-CO39 and AVR-Pia are founders of a family of sequence-unrelated but structurally conserved fungal effectors in a broad range of ascomycete phytopathogens (de Guillen et al., 2015). Consistent with structural modelling (IntFOLD Version 3.0 (McGuffin et al., 2015)), the recently resolved NMR-based and crystal structure of Bgh CSEP0064 revealed a ribonuclease-like fold (Pennington et al., 2019) with the absence of canonical catalytic residues in the substrate-binding pocket, and the gene products of ~120 additional Bgh CSEPs very likely adopt a similar structure (Pennington et al., 2019). Structural similarity searches (IntFOLD Version 3.0) also suggested a ribonuclease-like fold for AVR_{A7} and AVR_{A13} (high and certain confidence at p=3.739E-3, and 6.174E-4, respectively), whereas no significant structural similarities were detected between CSEP0064 and AVR_{A1}, AVR_{A9}, AVR_{A10}, or AVR_{A22} (low and medium confidence at p>0.01). Instead, we find that AVR_{A9} may adopt a structural fold that is similar to an antimicrobial peptide called microplusin (p=6.014E-3). In addition, no obvious structural similarities were predicted between AVR_{A1}, AVR_{A9}, and AVR_{A10}/AVR_{A22}, suggesting that allelic MLA receptors are capable of detecting the presence of structurally unrelated Bgh effectors. This assumption is consistent with the recent finding that the wheat Mla ortholog Sr50 directly binds to the Basidiomycete stem rust (Pgt) avirulence effector AvrSr50 (Chen et al., 2017). This Basidiomycete effector most likely evolved independently from the Ascomycete Bgh effectors and lacks significant sequence and predicted structural similarity with the known AVRa effectors. We speculate that MLA receptors might have an exceptional propensity to directly detect unrelated pathogen effectors and that this feature might have facilitated the functional diversification of the receptor in the host population.

In a whole leaf context, race-specific disease resistance specified by MLA receptors to Bgh is invariably linked to the activation of localized host cell death (Boyd et al., 1995). NLR-mediated cell death likely contributes to the termination of biotrophic fungal pathogenesis, including that of powdery mildews, because this class of pathogens feeds on living plant cells. A striking feature of the functional diversification at Mla is the enormous variation in microscopic and macroscopic resistanceassociated Bgh infection phenotypes as shown with barley NILs carrying different Mla resistance specificities (Boyd et al., 1995). For instance, the onset of detectable host cell death can vary dramatically and can be both rapid and limited to the first attacked leaf epidermal cell, terminating early fungal pathogenesis, or can occur at later stages of fungal pathogenesis and involve numerous leaf mesophyll cells that subtend Bgh-infected epidermal cells (Boyd et al., 1995). Here we have employed co-transfection experiments of barley leaf protoplasts with Mla-AVRa pairs and protoplast cell death as a proxy for receptor activation, excluding the possibility that additional Bgh-derived molecules associated with Pattern-triggered immunity influence the timing of immune receptormediated cell death in this system. Although based on overexpression data, the significant variation in cell death phenotypes reported here could partly reflect variable Bgh infection phenotypes on different MLA NILs (Supplementary file 1, Boyd et al., 1995). In turn, these differences of infection phenotypes are possibly due to variations in the steady-state levels of the MLA receptors during Bgh infection, timing of Bgh-mediated AVR $_{\Delta}$ secretion and/or AVR $_{\Delta}$ steady-state levels in planta, or MLA-AVR_A pair-dependent receptor binding affinities. Whilst etablishing the relevance of the the latter requires future biochemical characterization of MLA-AVRA complexes, our work revealed a very strong binding of AVR_{A13}-V2 to MLA13 both in the split-LUC and yeast two-hybrid experiments (Figure 5), thereby uncoupling AVRA binding to the receptor from receptor activation, that is immune receptor-mediated cell death activation. Future biochemical and genetic experiments will clarify



whether the naturally occurring AVR_{A13} -V2 effector variant acts as a dominant negative ligand when co-expressed with the AVR_{A13} avirulence effector.

Materials and methods

Key resources table

Reagent type (species) or resource	Designation	Source or reference	Identifiers	Additional information
Strain (Blumeria graminis f. sp. hordei)	CC107, CC148, CC1, CC52, CC66, CC88, NCI, 63.5, A6, B103, Aby, Art, Will, OU14, RACE1, K1	Lu et al. (2016) doi:10.1073/pnas. 1612947113.	GEO:GSE83237	
Strain (Blumeria graminis f. sp. hordei)	DH14	Frantzeskakis et al. (2018) doi:10.1186/s 12864-018-4750-6.	GEO:GSE106282	
Strain (Blumeria graminis f. sp. hordeí)	K2, K3, K4, S11, S15, S16, S19, S20, S21, S22, S23. S25, S26	this paper	GEO:GSE110266	collected in 2017 on cv. Meridian and Keeper barley at the Max Planck Institute for Plant Breeding Research, Cologne, Germany (GPS 5°57′N, 6°51′E 5)
Recombinant DNA reagent	pIPKb002	Himmelbach et al. (2007) doi:10.1104/pp.107.111575.	NCBI:EU161568.1	pZmUBQ:GW, Spc ^R
Recombinant DNA reagent	pGWB517	Nakagawa et al. (2007) doi:10.1263/jbb.104.34.	NCBI:AB294484.1	p35S:GW-4Myc, Spc ^R
Recombinant DNA reagent	pXCSG-GW-mYFP	García et al. (2010) doi:10.1371/journal .ppat.1000970.	NA	p35S:GW-mYFP, Carb ^R
Recombinant DNA reagent	pB42AD-GW	Shen et al. (2007) doi:10.1126/science. 1136372.	NA	pGal1:B42-AD- –1xHA-GW, TRP
Recombinant DNA reagent	pLexA-GW	Shen et al. (2007) doi:10.1126/science. 1136372.	NA	pADH1:LexA- BD-GW, HIS3
Recombinant DNA reagent	pDest-GW-nLUC	Gehl et al. (2011) doi:10.1111/j.1365- 313X.2011.04607.x.	NA	p35S:GW-Nterminus Luciferase, Kan ^R
Recombinant DNA reagent	pDest-GW-cLUC	Gehl et al. (2011) doi:10.1111/j.1365- 313X.2011.04607.x.	NA	p35S:GW-Cterminus Luciferase, Kan ^R
Gene (Blumeria graminis f. sp. hordei)	AVR _{a7} variants	Frantzeskakis et al. (2018) doi:10.1186/s12864- 018-4750-6.	csep0059; BLGH_06689; BLGH_06672; BGHR1_17217; BGHR1_17236; BGHR1_17237	
Gene (Blumeria graminis f. sp. hordeî)	AVR _{a9} variants	Frantzeskakis et al. (2018) doi:10.1186/s 12864-018-4750-6.	csep0174; BLGH_04994; BGHR1_10042	
Gene (Blumeria graminis f. sp. hordei)	AVR _{a10} /AVRa _{a22} variants	Frantzeskakis et al. (2018) doi:10.1186/s 12864-018-4750-6	csep0141; BLGH_05021; BGHR1_10013	

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Continued

Reagent type (species) or resource	Designation	Source or reference	Identifiers	Additional information
Gene (Blumeria graminis f. sp. hordei)	AVR _{a1} variants	Lu et al. (2016) doi:10.1073/ pnas.1612947113; Frantzeskakis et al. (2018) doi:10.1186/s 12864-018-4750-6	csep0008; BLGH_03023; BLGH_03022; BGHR1_11142	
Gene (Blumeria graminis f. sp. hordei)	AVR _{a13} variants	Lu et al. (2016) doi:10.1073/pnas.1612947113; Frantzeskakis et al. (2018) doi:10.1186/s 12864-018-4750-6	csep0372; BLGH_02099; BGHR1_12484	
Gene (Hordeum vulgare)	Mla9	Seeholzer et al. (2010) doi.org/10.1094 /MPMI-23-4-0497	NCBI: GU245941.1	
Gene (Hordeum vulgare)	Mla22	Seeholzer et al. (2010) doi.org/10.1094/ MPMI-23-4-0497	NCBI:GU245946	
Gene (Hordeum vulgare)	Mla10	Seeholzer et al. (2010) doi.org/10.1094/ MPMI-23-4-0497	Mla10	Different from NCBI:AY266445.1
Gene (Hordeum vulgare)	Mla7	Seeholzer et al. (2010) doi.org/10.1094/ MPMI-23-4-0497; Lu et al. (2016) doi:10.1186/s 12864-018-4750-6	Mla7 Different from NCBI:AY26644	
Gene (Hordeum vulgare)	Mla7 (AAQ55540_ Halterman et al., 2004)	Halterman and Wise (2004) doi:10.1111/j. 1365-313X.2004.02032.x	NCBI:AY266444.1	
Gene (Hordeum /ulgare)	Mla1	Seeholzer et al. (2010) doi.org/10.1094/ MPMI-23-4-0497; Lu et al. (2016)	NCBI:GU245961	
Gene (Hordeum vulgare)	Mla13	Seeholzer et al. (2010) , Lu et al. (2016) doi:10.1073/ pnas.1612947113.	AF523678.1	
Antibody	monoclonal rat anti-HA	Merck	3F10, RRID:AB_390914	1:2000
Antibody	monoclonal mouse anti-LexA	Santa Cruz Biotechnology	sc7544, RRID:AB_627883	1:1000
Antibody	polyclonal rabbit anti-c-myc	Abcam	ab9106, RRID:AB_307014	1:5000
Antibody	polyclonal rabbit anti-GFP	Abcam	ab6556, RRID:AB_305564	1:5000
Antibody	polyclonal rabbit anti-LUC	Sigma	L0159, RRID:AB_260379	1:2000
Antibody	polyclonal goat anti-rat IgG-HRP	Santa Cruz Biotechnology	sc2065, RRID:AB_631756	1:100 000
Antibody	polyclonal goat anti-mouse IgG-HRP	Santa Cruz Biotechnology	sc2005, RRID:AB_631736	1:100 000
Antibody	polyclonal donkey anti-rabbit IgG-HRP	Santa Cruz Biotechnology	sc-2313, 1:100 000 RRID:AB_641181	
Antibody	monoclonal rabbit anti-GFP	Santa Cruz Biotechnology	sc-8334, RRID:AB_641123	1:5000



Plant materials and growth conditions

The barley cultivars (cv.) Golden Promise, Manchuria, and Pallas and their near isogenic lines (*Kolster et al., 1986*; *Moseman, 1972*), were grown at 19°C, 70% relative humidity, and under a 16 h photoperiod. *Nicotiana benthamina* plants were grown in standard greenhouse conditions under a 16 h photoperiod.

Fungal isolates

Barley leaves suspected of being infected by *Bgh* were collected from the cv. Meridian and Keeper barley fields at the Max Planck Institute for Plant Breeding Research, Cologne, Germany (GPS 5°57′N, 6°51′E 5). Spores of different infected field leaves were transferred onto one-week-old barley leaves of the cv. Manchuria (lacking any *Mla* resistance specificity). Inoculated Manchuria leaves were incubated on 1% Bacto Agar plates supplemented with 1 mM benzimidazole at 20°C, 70% humidity, and long-day conditions for one week until *Bgh* conidiospore growth was visible. Subsequent single spore propagation was applied three (S11, S15, S19, S20, S21, S23, S25, S26) or six (K2, K3, K4, S16 and S22) times for isolation of single *Bgh* isolate genotypes. In total, we collected 13 *Bgh* isolates, which were tested at least three times on a panel of cv. Pallas and cv. Manchuria *Mla* near-isogenic lines. Maintenance of fungal isolates and other *Bgh* isolates in this study was carried out as described previously (*Lu et al., 2016*).

RNA sequencing

The new RNA-seq data generated for this study are deposited in the National Center for Biotechnology Information Gene Expression Omnibus (GEO) database (accession no. GSE110266). The previously generated RNA-seq data for DH14 and all other isolates can also be found at GEO (accession nos. GSE106282 and GSE83237, respectively).

RNA-seq read alignment and variant calling

The RNA-seq reads for all datasets were mapped to the new *Bgh* DH14 reference genome assembly (version 4.0) taking into account exon-intron structures using the splice-aware aligner Tophat2 (*Kim et al., 2013*), which considers known splice sites based on the new DH14 gene models (*Frantzeskakis et al., 2018*). Read length was 100 bp for previously sequenced isolates (GSE83237), 150 bp for DH14 (GSE106282), S20 and S25, and 250 bp for all other isolates (GSE110266). To allow for adequate alignment efficiency also for those isolates with higher sequence divergence from the reference genome, we adjusted the alignment settings as follows: –read-mismatches 10 –read-gaplength 10 –read-edit-dist 20 –read-realign-edit-dist 0 –mate-inner-dist 260 –mate-std-dev 260 –minanchor 5 –splice-mismatches 2 –min-intron-length 30 –max-intron-length 10000 –max-insertion-length 20 –max-deletion-length 20 –num-threads 10 –max-multihits 10 –coverage-search –library-type fr-firststrand –segment-mismatches 3 –min-segment-intron 30 –max-segment-intron 10000 – min-coverage-intron 30 –max-coverage-intron 10000 –b2-very-sensitive. Using the SAMtools suite (Version 0.1.18) (*Li et al., 2009*), the generated alignment files were subsequently filtered to retain only properly paired reads with mapping quality >0 for the downstream analyses.

To assess the expression levels of individual genes, we obtained the fragment counts per gene for each isolate and time point from the mapped RNA-seq reads after filtering using the Subread function 'featureCounts' (version 1.5.0) (*Liao et al., 2014*) with options -t CDS -s 2 M -p. Subsequently, the raw counts were summarized over both time-points for each isolate and normalized to FPKM (fragments per kilobase of transcript per million mapped reads) values for better comparability of expression levels.

In parallel, sequence variants were identified from the mapped RNA-seq reads using two different tools. In both cases, the variant calling was performed on a combined alignment dataset that was obtained by merging the mapped RNA-seq reads from all isolates using the merge function of the SAMtools suite (*Li et al., 2009*). In one approach, single nucleotide polymorphisms (SNPs) were identified using the mpileup function in the SAMtools suite (*Li et al., 2009*) with options -A, -u, -D, -d 30000 -L 7000. The resulting mpileup variants were filtered using SnpSift (Version 3.4) (*Cingolani et al., 2012a*) with filter settings "(AF1 >= 0.01852) and ((DP >= 30) | ((DP >= 10) and (GEN[*].DP>=5))) and (QUAL >= 50) and (GEN[*].GQ>=10) and ((na PV4) | ((PV4[0]>1e-10) and (PV4 [3]>1e-5)))" to extract high-quality variants with sufficiently high allele frequency (\geq 0.01852; that is



alternate allele present in at least ~50% of the reads of one isolate), sufficient read coverage (≥30 reads in total, or \geq 10 reads in total and at least one isolate with \geq 5 reads), a SNP calling quality score >50, at least one isolate with a genotype quality score >10, and absence of extreme placement bias. In the other approach, variants were called using freebayes (version 9.9.2) (Garrison and Marth, 2012) with options -ploidy 1 -use-duplicate-reads -min-mapping-quality 0 -min-base-quality 20 -min-coverage 30 -genotype-qualities. To allow correct variant calling from our RNA-seg data with freebayes, the mapped reads in this case were preprocessed using the function SplitNCigar-Reads in the GenomeAnalysis Toolkit (version 3.4.0) (McKenna et al., 2010) with options -U ALLOW_N_CIGAR_READS -fixNDN -maxOverhang 10 to split any reads with splice junctions. An additional, independent freebayes variant calling was also performed with -ploidy 2, to allow processing of cases where an isolate contains additional gene copies that differ from each other at some residue(s). The resulting freebayes variant sets were filtered using SnpSift (Cingolani et al., 2012a) with filter settings "(AF[*]>=0.01852) and ((exists GEN[*].GQ) and (GEN[*].GQ[*]>=10)) and ((exists GEN[*].AO) and (GEN[*].AO[*]>=3)) and ((DP >= 30) | ((DP >= 10) and (exists GEN[*].DP) and (GEN [*].DP>=5))) and (SAR[*]>0) and (SAF[*]>0) and (RPR[*]>1) and (RPL[*]>1)' to extract high-quality variants fulfilling the same criteria as described above for mpileup. The subsequent variant annotation and effect prediction for all datasets was performed using snpEff (Version 3.4; default settings) (Cingolani et al., 2012b) based on the new DH14 genome and gene models.

Population structure and genetic association analysis of Bgh isolates

To obtain a suitable set of single nucleotide polymorphisms (SNPs) for population structure analysis, we further filtered the set of high-quality variants obtained with mpileup from the combined alignment data as described above. We extracted only silent (synonymous) SNPs in coding regions for which exactly two different alleles were found (diallelic SNPs), and complete genotype information was available for all isolates (i.e., no missing data). The resulting set of 6286 high-quality diallelic synonymous SNPs was used to examine the genotype data for the presence of any obvious population structure using the R packages adegenet (Version 2.0.1) (*Jombart and Ahmed, 2011*) and ape (Version 3.4) (*Paradis et al., 2004*). To this end, we created a PCA plot from the genotype data using the function glPca (R package adegenet) and additionally computed a neighbour-joining tree based on the pairwise Euclidean distances between the isolate genotypes using the function nj (R package ape). Additionally, another PCA plot was generated for the European isolates only, based on a set of 5170 high-quality diallelic synonymous SNPs found in the European isolates.

For the association analysis we focused on the high-quality variants obtained with freebayes in the haploid SNP calling, which we filtered further to extract only non-synonymous coding variants predicted to change the protein sequence and generated a simplified genotyping table listing all of these variants. Additionally, we also screened the results from the diploid freebayes variant calling for 'heterozygous' positions with a minor allele frequency of at least 1/3, as in this case the different 'alleles' are likely derived from differing paralog copies, and added these positions to the genotyping table. This procedure was implemented in R (*Supplementary file 2*) and resulted in a set of 22,838 high-confidence variants with predicted effect on the protein sequence, which we tested for their association with the observed avirulence phenotypes using Fisher's exact test.

Loss of avirulence might be caused by different variants in different isolates. Therefore, we integrated all high-confidence non-synonymous variants over each gene to obtain gene-wise genotypes. Moreover, to also include presence/absence polymorphisms we considered the complete absence of a transcript as a 'missing' genotype. Finally, these gene-wise genotypes were tested for association with the observed avirulence phenotypes using Fisher's exact test. This gene-wise integration of variants and the subsequent association test were implemented in R. As further technical validation, we additionally performed the same association test also on gene-wise genotypes obtained in the same way from the high-quality mpileup SNPs (*Supplementary file 3*). All identified *AVR*_a candidates were picked up using both tools; the p values mentioned are based on the freebayes variants (*Supplementary file 4*).

Genome sequencing, assembly and annotation

For the improvement of the *Bgh* isolate K1 assembly, genomic DNA was extracted as described (*Feehan et al., 2017*) and sequenced using the Oxford Nanopore MinION platform according to the



manufacturer's instructions for library generation and flow-cell handling. Base-Calling of the resulting long-reads was performed with the Albacore Sequencing Pipeline Software (version 2.0.2, Oxford Nanopore) yielding 3,09 GB of data in 781831 reads, and subsequently assembled with Canu (v1.4, Koren et al., 2017). The 2238 assembled contigs were corrected using Illumina short reads (SRR650349 and SRR654727; Hacquard et al., 2013) with Pilon (v1.18, Walker et al., 2014) in four iterations. Re-annotation of the new genome assembly was performed as described in Frantzeskakis et. al., and CSEPs were then manually curated using WebApollo (v2.0.6, Lee et al., 2013). Data are deposited under the accession number PRJEB30373 at EBI-ENA.

Genome-wide nucleotide diversity of B.g. ff. spp. isolates

Illumina short-read sequences of different *B.g.* ff. ssp. (SRA accession number: SRP062198, *Supplementary file 5*) were mapped to the reference genome of isolate 96224 (t ENA accession number: PRJEB28180) as described (*Müller et al., 2019*). For extraction of neutral markers, SNP calling was done with freebayes with default parameters (*Garrison and Marth, 2012*) and SNPs located within genes were excluded (*Müller et al., 2019*). Vcftools (*Danecek et al., 2011*) was used to filter SNPs with the following options: vcf –remove-indels –max-alleles 2 –min-alleles 2 –minDP 8 –maxDP 100 –max-missing 1 –recode –maf 0.01 –minGQ 20. Per-site nucleotide diversity was calculated with vcftools –sites-pi command.

Maximum likelihood phylogeny for predicted secreted proteins

In order to generate a maximum-likelihood phylogenetic tree based on the mature peptide sequences of the Bgh DH14 SPs, the 805 predicted secreted protein sequences were aligned using MAFFT v7.310 (*Katoh and Standley, 2013*) with the settings –maxiterate 1000 –localpair. The alignment was then passed to IQTree v1.6.beta4 (*Nguyen et al., 2015*) with the settings -nt 10 -mem 12G -bb 1000. The resulting tree was then visualized using iTOL (*Letunic and Bork, 2016*).

Generation of expression constructs

All genes with or without stop codons, were amplified from the cDNA of Bgh isolates (Lu et al., 2016) or plasmid templates (Seeholzer et al., 2010) using Phusion Hot Start II high-fidelity DNA polymerase (Thermo Scientific) and subsequently cloned into pENTR/D-TOPO (Km^R) (Thermo Scientific) or synthesized as pDONR221 (Km^R) entry clones from GeneArt (Thermo Scientific). The sequence integrity of all clones was confirmed by Sanger sequencing (Eurofins). Primers for AVR_a amplification were designed to replace the signal peptide with the ATG start codon (Supplementary file 6).

For transient gene expression assays in planta and for yeast 2-hybrid interaction studies, respective genes were transferred from entry or donor vectors into the expression vectors pIPKb002 (Spc^R) (Himmelbach et al., 2007), pGWB517 (Spc^R) (Nakagawa et al., 2007), pXCSG-GW-mYFP ($Carb^R$) (García et al., 2010), pLexA-GW ($Carb^R$), or pB42AD-GW ($Carb^R$) (Shen et al., 2007) as indicated using LR Clonase II (Thermo Scientific).

For the split-LUC assay, genes of interest were transferred from expression vectors into pDONR207 (Gm^R) using BP clonase II (Thermo Scientific) and subsequently cloned into pDEST-GW-nLUC (Km^R) or pDEST-GW-cLUC (Km^R) ($Gehl\ et\ al.$, 2011) using LR Clonase II. Alternatively, pENTR/D-TOPO or pDONR221 entry clones were double-digested using Pvul and Nrul (NEB) to remove Km^R and linearized constructs were transferred directly into pDEST-GW-nLUC (Km^R) or pDEST-GW-cLUC (Km^R) using LR clonase II; the integrity of the resulting expression constructs was examined by Sanger sequencing (Eurofins).

Transient gene expression and cell death assay in barley protoplasts

Assessment of protoplast cell death using a luciferase activity as a proxy for cell viability was adapted from ($Lu\ et\ al.$, 2016) with the following modifications: cDNAs of the AVR_a candidate genes lacking their respective signal peptides (SPs) were co-expressed together with cDNAs of the corresponding MLA receptors from the same strong ubiquitin promotor in the same barley genetic background to directly compare cell death activities mediated by different MLA and AVR_A pairs. For this, the epidermis of the second leaves from seven to eight-day-old plants of the cultivar Golden Promise was removed before leaves were immersed in the enzyme solution. A total volume of 35 μ l water



containing 5 μ g of the *luciferase* reporter plasmid, 12 μ g of the respective *Mla* construct, and 6 μ g of the respective effector construct or an EV was transfected into 300 μ L barley protoplasts at a concentration of 3 \times 10⁵ protoplasts/ml solution. For co-transfection of AVR_{a10} and EKA_AVR_{a10} , 5 μ g of the *luciferase* reporter plasmid, 10 μ g of the *Mla10* construct, 4 μ g of AVR_{a10} , and either 4 μ g of EKA_AVR_{a10} or 4 μ g of EV were transfected. At 16 hr after transfection, protoplasts were collected by centrifugation at 1000 \times g, the supernatant was discarded, and 200 μ l 2x cell culture lysis buffer were added (Promega, E1531). Luciferase activity was determined by mixing 50 μ l of protoplast lysate with 50 μ l luciferase substrate (Promega, E1501) in a white 96-well plate and light emission was measured 1 s/well using a microplate luminometer (Centro, LB960). Relative luciferase reads (*Figure 4—source datas 1–3*, *Figure 4—figure supplement 1—source datas 1–3*), were calculated by setting the control empty vector sample read of each individual experiment to 1.

Transient gene expression by Agrobacterium-mediated transformation of Nicotiana benthamiana leaves

Agrobacterium tumefaciens GV3101::pMP90 and A. tumefaciens GV3101::pMP90K were freshly transformed with respective constructs of interest and grown from single colonies in liquid Luria broth medium containing appropriate antibiotics for ~24 hr at 28°C to an OD₆₀₀ not higher than 1.5. Bacterial cells were harvested by centrifugation at 2500 \times g for 15 min followed by resuspension in infiltration medium (10 mM MES, pH 5.6, 10 mM MgCl₂, and 200 μ M acetosyringone) to a final OD₆₀₀ = 1.2. Cultures were incubated for 2 to 4 hr at 28°C with 180 rpm shaking before infiltration into leaves from three to five-week-old N. benthamiana plants. Bacteria carrying AVR_a constructs or EV plasmid were mixed equally with Mla plasmid-carrying bacteria. Tissue for immunodetection analysis was harvested two days post infiltration and cell death scores (Figure 4—figure supplement 2—source datas 4,5) were assessed three days post infiltration throughout.

Plant protein extraction and pull-down for fusion protein detection by immunoblotting

Frozen leaf material was ground to a fine powder using pre-cooled adapters in a bead beater (Retsch) and thawed in cold plant protein extraction buffer (150 mM Tris-HCl, pH 7.5, 150 mM NaCl, 10 mM EDTA, 10% (v/v) glycerol, 5 mM DTT, 2% (v/v) plant protease inhibitor cocktail (Sigma), 1 mM NaF, 1 mM Na $_3$ VO $_4$, 1 mM PMSF, and 0.5% (v/v) IGEPAL) at a ratio of 150 mg fresh tissue/1 ml of extraction buffer. Extracts were centrifuged twice at 15,000 \times g for 15 min at 4°C. For SDS-PAGE, extracts were diluted 4:1 with 4x SDS loading buffer and heated to 95°C for 5 min.

For pull-down of mYFP-tagged proteins, GFP-Trap-MA (Chromotek) beads were incubated in equilibration buffer (*Saur et al., 2015*) for 1 hr at 4°C and subsequently mixed with protein extracts for 2 to 3 hr at 4°C with slow but constant rotation. Then, conjugated GFP-Trap beads were washed five times in 1 ml of cold wash buffer (*Saur et al., 2015*) at 4°C before interacting proteins were stripped from the beads by boiling in 25 μ l of 4x SDS loading buffer for 5 min.

Samples were separated on 10% SDS-PAGE gels, blotted onto PVDF membrane, and probed with anti-GFP (Santa Cruz Biotechnology sc-8334, RRID:AB_641123; or abcam ab6556, RRID:AB_305564), anti-LUC (Sigma L0159) or anti-c-Myc (abcam ab9106, RRID:AB_307014), followed by anti-rabbit IgG-HRP (Santa Cruz Biotechnology sc-2313, RRID:AB_641181) secondary antibodies. Proteins were detected by the HRP activity on SuperSignal West Femto Maximum Sensitivity Substrate (Thermo Fisher 34095) using a Gel Doc XR +Gel Documentation System (Bio-Rad).

Split-luciferase complementation assay

To obtain protein extracts for luciferase measurements, one leaf disk with a diameter of 0.38 cm was harvested from three different leaves at two days post transformation, resulting in three leaf disks/sample. Samples were frozen in liquid nitrogen and ground to a fine powder using a pre-cooled adapter in a Retsch bead beater. Each sample was thawed in 100 μ l 2x cell culture lysis buffer (Promega, E1531) supplemented with Tris-HCl, pH 7.5 to a final concentration of 150 mM. Luciferase activity was determined by mixing 50 μ l of leaf extract with 50 μ l luciferase substrate (Promega, E1501) in a white 96-well plate and light emission (*Figure 5—source datas 1–3*) was measured 1 s/well in a microplate luminometer (Centro, LB960). Complementation of a functional LUC protein by genetic fusion of bait/prey with the nucleotides encoding the N-terminal 416 aa of LUC (nLUC) and



C-terminal 152 aa of LUC (cLUC) allows the detection of a real-time and reversible signal for direct interaction (*Bosmans et al., 2016; Chen et al., 2008*).

Yeast 2-hybrid assay and yeast protein extraction

Mla variants were cloned into the pLexA-GW vector (*Shen et al., 2007*) for expression with an N-terminal LexA activation domain under the control of a constitutive ADH1 promoter (BD-MLA). The AVR_a variants were cloned into pB42AD-GW (*Shen et al., 2007*) for expression with an N-terminal B42 activation domain followed by the HA-tag under the control of an inducible GAL1 promoter (AD-AVR_A). Using the lithium acetate method (*Gietz and Woods, 2002*), Mla bait constructs and AVR_{a13} prey constructs were co-transformed into the yeast strain EGY4.8 p8op-lacZ and successful transformants were selected by colony growth on SD-UHW/Glu (4% (w/v) Glucose, 0.139% (w/v) yeast synthetic drop-out medium pH 5.8 without uracil, histidine, tryptophan, 0.67% (w/v) BD Difco yeast nitrogen base, 2% (w/v) Bacto Agar). Yeast transformants were grown to $OD_{600} = 1$ in liquid SD-UHW/Glu before harvesting cells for drop out of the dilution series on SD-UHW/Gal/Raf media (SD-UHW without glucose but with 2% (w/v) Galactose 1% (w/v) Raffinose, with (-UHW) or without Leucine (-UHWL)) and incubated for six days at 30°C followed by room temperature incubation for another six days.

For protein detection, yeast strains were grown to $OD_{600} = 1$ in SD-UHW/Gal/Raf liquid medium at 30°C and 200 rpm shaking, and proteins were extracted using 200 mM NaOH (NaOH method; **Zhang et al., 2011**). Total protein samples were separated on 9% or 12% SDS-PAGE gels, blotted onto PVDF membrane, and probed with anti-HA (Merck, clone 3F10, RRID:AB_390914) or anti-LexA (Santa Cruz Biotechnology, sc7544, RRID:AB_627883) primary antibodies followed by anti-rat (Santa Cruz Biotechnology, sc2065, RRID:AB_631756) or anti-mouse IgG-HRP (Santa Cruz Biotechnology, sc2005, RRID:AB_631736) secondary antibodies as appropriate. HA and LexA fusion proteins were detected by HRP activity on SuperSignal West Femto Maximum Sensitivity Substrate (Thermo Fisher 34095) using a Gel Doc XR +Gel Documentation System (Bio-Rad).

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Additional files

Supplementary files

• Supplementary file 1. Infection phenotypes of *Bgh* isolates on the Pallas and Manchuria cultivar accessions used for the association test.

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• Supplementary file 2. Script for freebayes genetic association analysis.

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• Supplementary file 3. Script for mpileup genetic association analysis.

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• Supplementary file 4. Statistical summary of association analysis.

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• Supplementary file 5. Blumeria graminis isolates used for phylogenetic analysis of CSEP0141 ($AVR_{a10}/AVRa_{a22}$).

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• Supplementary file 6. Primers used in this study.

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• Transparent reporting form

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Data availability

RNA sequencing data have been deposited in GEO under accession code GSE110266 and improved Blumeria graminis f.sp. hordei isolate K1 assembly is deposited under the accession number PRJEB30373 at EBI-ENA. All data generated or analysed during this study are included in the manuscript and supporting files.

The following datasets were generated:

Database and



Author(s)	Year	Dataset title	Dataset URL	Identifier
Isabel ML Saur, Sas- kia Bauer, Barbara Kracher, Lamprinos Franzeskakis, Mar- ion C Müller, Björn Sabelleck, Florian Kümmel, Ralph Panstruga, Paul Schulze-Lefert	2019	Improved Blumeria graminis f.sp. hordei isolate K1 assembly	https://www.ebi.ac.uk/ ena/data/search?query= PRJEB30373	European Nucleotide Archive, PRJEB30373
Saur IML, Bauer S, Kracher B, Lu X, Franzeskakis L, Müller MC, Sabel- lack B, Kümmel F, Panstruga R, Mae- kawa T, Schulze- Lefert P	2018	Identification of the AVRa7,AVRa9, AVRa10 and AVRa22 effector genes from barley powdery mildew fungus (Bgh) association analysis between transcript polymorphisms and AVRa phenotypes from 27 Bgh isolates	nih.gov/geo/query/acc.	NCBI Gene Expression Omnibus, GSE110266

The following previously published dataset was used:

Author(s)	Year	Dataset title	Dataset URL	Database and Identifier
Lu X, Kracher B, Saur IML, Bauer S, Ellwood S, Wise R, Yaeno T, Maekawa T	2016	Identification of avirulence genes in the barley powdery mildew fungus (Bgh) by RNA-sequencing and transcriptome-wide association analysis on a set of Bgh isolates		NCBI Gene Expression Omnibus, GSE83237

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