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Host-specificity factors in plant pathogenic fungi

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ABSTRACT

Fortunately, no fungus can cause disease on all plant species, and although some plant-pathogenic fungi have quite a broad host range, most are highly limited in the range of plant species or even cultivars that they cause disease in. The mechanisms of host specificity have been extensively studied in many plant-pathogenic fungi, especially in fungal pathogens causing disease on economically important crops. Specifically, genes involved in host specificity have been identified during the last few decades. In this overview, we describe and discuss these host-specificity genes. These genes encode avirulence (Avr) proteins, proteinaceous host-specific toxins or secondary metabolites. We discuss the genomic context of these genes, their expression, polymorphism, horizontal transfer and involvement in pathogenesis.

1. Introduction

Plant diseases caused by fungi are recognized as a major threat to food security (Doehlemann et al., 2017; Fisher et al., 2012; Pennisi, 2010). For example, wheat stem rust caused by Puccinia graminis, Asian soybean rust caused by Phakopsora pachyrhizi, rice blast caused by Magnaporthe oryzae, and banana black sigatoka caused by Mycosphaerella fijiensis have resulted in serious yield losses in human history (Pennisi, 2010).

Most plant pathogenic fungal species have a narrow range of plant species in which they cause disease, a phenomenon we here call ‘host species specificity’. The collective host range of some fungal species, such as Fusarium oxysporum (F. oxysporum), can be very large, but then individual strains are often limited to infect one or a few plant species only (Borah et al., 2018; Pietro et al., 2003). Based on host range, strains within such fungal species are commonly classified into different pathotypes or formae speciales. In addition, fungal species or formae speciales are sometimes divided into different races depending on the particular cultivars of a plant species that they are able to infect. This phenomenon we here call ‘host cultivar specificity’. For example, F. oxysporum is classified into more than 100 formae speciales based on host species specificity, including the tomato-infecting strain F. oxysporum forma specialis (f. sp.) lycopersici (Fol), while Fol itself is subdivided into three races based on host cultivar specificity (Takken and Rep, 2010). The molecular basis of host cultivar specificity has been extensively studied in many plant pathogenic fungi, while the molecular basis of host species specificity is less well understood (de Wit, 2016; Lanver et al., 2017; Lo Presti et al., 2015; Prasad et al., 2019; Selin et al., 2016; Yan and Talbot, 2016). In this review, the term ‘host specificity’ includes both host species specificity and host cultivar specificity.

To understand the genetic basis of host specificity in plant pathogenic fungi, the emergence of a molecular understanding of plant immunity over the last three decades has been essential. To defend themselves against fungal pathogens, plants have evolved two layers of immunity (Jones and Dangl, 2006). The first layer of immunity responds to pathogen-associated molecular patterns (PAMPs) common to many microbes, including non-pathogens, and this defense system is called PAMP-triggered immunity (PTI) (Jones and Dangl, 2006). To suppress PTI responses, pathogens secrete molecules called effectors to facilitate colonization. These effectors are commonly secreted proteins but can also be metabolites. Some effectors target plant susceptibility (S) proteins, resulting in effector-triggered susceptibility (ETS) (van Schie and Takken, 2014). For example, necrotrophic effectors of Stagonospora nodorum (S. nodorum) are able to interact with wheat susceptibility gene products (Oliver et al., 2012). The second layer of defense comprises plant resistance (R) proteins that directly or indirectly recognize pathogen-produced effectors, resulting in effector-triggered immunity (ETI) (Jones and Dangl, 2006). Among identified R proteins, most are nucleotide-binding leucine-rich repeat proteins (NLRs) (Kourelis and van der Hoorn, 2018). To evade effector recognition by plant R proteins, pathogens can undergo loss or mutation of the corresponding effector genes (Jones and Dangl, 2006; Wang and Wang, 2018). As a result, pathogens and plants are evolving in a perpetual arms race. Most effector genes are located in repeat-rich regions. For example, all known effectors genes in Leptosphaeria maculans (L.}
2.1.1. F. oxysporum f. sp. lycopersici (Fol)
The I-3 protein, was identified in Fol as well (Rep et al., 2004). Resistance gene recognition by I-2, resulting in race 3 (Houterman et al., 2009). AVR2 (Houterman et al., 2009; Ma et al., 2015). All three Fol AVR genes encode small, secreted proteins with multiple cysteines, and are located on a single accessory chromosome with high density of repetitive elements (Ma et al., 2010; Schmidt et al., 2013).

This arms race between pathogens and plants results in relatively fast evolution of effectors, R and S proteins (Jones and Dangl, 2006; Wang and Wang, 2018). In general, effectors promote virulence and are therefore virulence factors. Effectors that are recognized by R proteins are (also) called avirulence (Avr) factors (Avr proteins) (Stotz et al., 2014). These avirulence and virulence factors determine resistance and susceptibility of plants, respectively, are considered to be host-specificity factors and have been identified in many plant pathogenic fungi (de Wit, 2016; Lanver et al., 2017; Selin et al., 2016; Toruño et al., 2016; Yan and Talbot, 2016). The virulence factors reviewed here are known as proteinaceous toxins, therefore they are referred as proteinaceous host-specific toxins hereafter. In addition to pathogen-secreted proteins that can determine host specificity, secondary metabolites can also act as host-specificity determinants, such as the host-selective toxins (HSTs) of Alternaria alternate and Verticillium dahliae (Chen et al., 2018; Tsuge et al., 2013; Zhang et al., 2019).

Here we first describe host-specificity factors in plant pathogenic fungi, including avirulence proteins, proteinaceous host-specific toxins and secondary metabolites. Then, involvement of horizontal transfer and the genomic context in the acquisition and evolution of host-specificity genes is addressed.

2. Avirulence proteins determining host specificity

Avirulence proteins determining host cultivar specificity have been identified in various fungal species (Table 1). Below we provide a brief overview of these cases, including involvement in pathogenesis, polymorphism and interaction with the corresponding resistance proteins.

2.1. Fusarium oxysporum

F. oxysporum is a soil-borne (presumably) asexual fungal species complex, which causes vascular wilt and root rot (Edel-Hermann and Lecomte, 2019). Avirulence genes in F. oxysporum determining host cultivar specificity have been identified in several formae speciales, including Fol (Takken and Rep, 2010), F. oxysporum f. sp. melonis (Schmidt et al., 2016) and F. oxysporum f. sp. niveum (Niu et al., 2016). Here, we will review the avirulence (AVR) genes identified in Fol and F. oxysporum f. sp. melonis, which infect tomato and melon, respectively.

2.1.1. F. oxysporum f. sp. lycopersici (Fol)

F. oxysporum f. sp. lycopersici (Fol) can be divided into three races based on their capability to infect tomato cultivars containing different resistance genes to Fol (Takken and Rep, 2010). Race 1 contains three AVR genes, notably AVR1, AVR2 and AVR3. The protein encoded by resistance gene I recognizes the product of AVR1, upon which the immune system is activated in the plant (Houterman et al., 2008). Avr1 also suppresses recognition of Avr2 and Avr3 by resistance proteins I-2 and I-3, respectively (Houterman et al., 2008). Race 2 evolved from race 1 by deletion of a chromosomal region containing AVR1, likely due to a recombination event between two TEs bordering the fragment (Biju et al., 2017). The I-2 resistance gene was introduced into tomato cultivars to protect them against race 2. The I-2 protein recognizes Avr2 (Houterman et al., 2009; Ma et al., 2015). Single point mutations in AVR2 subsequently emerged such that the gene product was no longer recognized by I-2, resulting in race 3 (Houterman et al., 2009). Resistance gene I-3 against race 3 was introduced in tomato cultivars, and the corresponding AVR3 gene, the product of which is recognized by the I-3 protein, was identified in Fol as well (Rep et al., 2004). Among the three Fol AVR genes known, AVR1 is not required for full virulence on susceptible hosts (Houterman et al., 2008), whereas AVR2 and AVR3 are (Houterman et al., 2009; Rep et al., 2004). For activation of I-2-mediated resistance, not only Avr2 is required, but also a Fol protein called Secreted in xylem 5 (Six5). Like Avr2, Six5 is required for full virulence (Houterman et al., 2009; Ma et al., 2015). All three Fol AVR genes encode small, secreted proteins with multiple cysteines, and are located on a single accessory chromosome with high density of repetitive elements (Ma et al., 2010; Schmidt et al., 2013).

2.1.2. F. oxysporum f. sp. melonis

F. oxysporum f. sp. melonis is divided into race 0, race 1, race 2 and race 1,2. So far, only one avirulence gene has been identified, AvrFom2, whose product is recognized by the protein encoded by the melon R gene Fom2 (Schmidt et al., 2016). AvrFom2 is a small secreted protein with two cysteine residues and without recognizable domains (Schmidt et al., 2016). However, it does show an overall low similarity to ToxA of Pyrenophora tritici-repentis (discussed below), and the cysteine residues that form the characteristic cysteine knot in ToxA are conserved (Schmidt et al., 2016). AvrFom2 is located in a lineage-specific region of the Fom001 genome and resides close to transposons (Schmidt et al., 2016; van Dam et al., 2017). The gene is absent in race 2 isolates (Schmidt et al., 2016).

2.2. Cladosporium fulvum

C. fulvum (Passalora fulva) is a non-obligate biotrophic fungal species and the causal agent of tomato leaf mold (de Wit, 2016). Already in the 1970s, it was found that the gene-for-gene relation between tomato and C. fulvum is based on the interaction of specific fungal products with specific resistance proteins in tomato (van Dijkman and Kaars Sijpsteijn, 1973).

The first fungal Avr protein was identified in C. fulvum in 1991 (van Kan, 1991). Until now, ten AVR genes of C. fulvum, which encode small cysteine-rich proteins, have been identified, notably Avr2, Avr4, Avr5 (Luderer et al., 2002), Avr4 (Joosten et al., 1994), Avr6 (van Dijkman and Kaars Sijpsteijn, 1973) and Avr9 (van Kan, 1991), Ecpl1 (Laugé et al., 1997), Ecpl2-1 (Laugé et al., 1997), Ecpl4 (Laugé et al., 2000), Ecpl5 (Laugé et al., 2000) and Ecpl6 (Bolton et al., 2008). Recognition of the encoded proteins in tomato is mediated by the cognate Cf resistance proteins Cf-2, Cf-4, Cf-4E, Cf-5 and Cf-9, Cf-Ecp1, Cf-Ecp2-2, Cf-Ecp4, Cf-Ecp5, Cf-Ecp6, respectively. These C. fulvum Avr proteins and another apoplastic effector proteins have been extensively reviewed previously (de Wit, 2016; Rivas and Thomas, 2005; Stergiopoulos and de Wit, 2009). In a recent study, nine newly identified small secreted proteins were found to be recognized by specific wild accessions of tomato, but the corresponding Cf immune receptor genes in these accessions are still unknown (Mesarič et al., 2018).

All C. fulvum Avr proteins identified are less than 300 amino acids in size and contain an even number of at least four cysteine residues (Mesarič et al., 2018, 2014). A virulence function for Avr2, Avr4, Avr5 and Ecpl6 has been demonstrated (Mesarič et al., 2014; Stergiopoulos and de Wit, 2009). To avoid recognition by R proteins, several types of sequence modifications have occurred in C. fulvum AVR genes including gene deletions, gene disruption by insertion of a transposon-like element, and nonsynonymous amino acid substitutions (Stergiopoulos and de Wit, 2009). The frequency of such mutations may have been enhanced by proximity of these AVR genes to repetitive elements (de Wit et al., 2012).

2.3. Leptosphaeria maculans

L. maculans is a hemi-biotrophic ascomycete responsible for stem canker of oilseed rape (Petit-Houdenet and Fudal, 2017). To date, eight AVR genes from L. maculans have been identified, Avrlm1, Avrlm2, Avrlm3, Avrlm4–7, Avrlm5–9 (AvrlmJ1), Avrlm6, Avrlm10 and Avrlm11, and all are located in repeat-rich, gene-poor genomic regions.
## Table 1
Host-specificity factors in plant pathogenic fungi.

| Fungal species | Host | Host-specificity genes | Corresponding resistance or susceptibility genes | Genome location of host-specificity genes | References |
|----------------|------|------------------------|--------------------------------------------------|-------------------------------------------|------------|
| *Fusarium oxysporum f. sp. lycopersici* | Tomato | AVR1, AVR2, AVR3 | I-1, I-2, I-3 | Accessory chromosome | (Ma et al., 2010; Takken and Rep, 2010) |
| *Fusarium oxysporum f. sp. melonis* | Melon | AvrFom2 | Fom2 | Close to repetitive elements | (Schmidt et al., 2016) |
| *Cladosporium fulvum* | Tomato | AvrLm1, AvrLm2, AvrLm3, AvrLm4-7, AvrLm5-9, AvrLm6, AvrLm10 and AvrLm11 | Rlm1, Rlm2, Rlm3, Rlm4, Rlm5, Rlm9, Rlm6, Rlm10, and Rlm11 | Repeat-rich regions | (Ghanbarian et al., 2018; Petit-Houdenot et al., 2019; Plissonneau et al., 2018; Rouxel and Balesdent, 2017) |
| *Magnaporthe oryzae* | Grass species | Av-P1, Av-P2, Av-P3, Av-P4, Av-P5, Av-P6, Av-P7, Av-P8, Av-P9, Av-P10, and Av-P11 | Pi-1, Pi-2, Pi-3, Pi-4, Pi-5, Pi-6, Pi-7, Pi-8, Pi-9, Pi-10, and Pi-11 | Close to repetitive elements | (Fernandez and Orth, 2018) |
| *Rhynchosporium secalis* | Barley | Np1 | Rs1 | Present on a large chromosome | (Hahn, 1993; Mohd-Assaad et al., 2019) |
| *Melampsora lini* | Flax | AvrLs67, AvrM, AvrP123, AvrP4, AvrL2A, and AvrM14-3 | L5, I6, L7, M, P, P1, P2, P3, P4, L14 | Mostly located at recombination hot-spots | (Lorrain et al., 2019; Petre et al., 2014) |
| *Puccinia graminis f. sp. tritici* | Wheat | AvrSs55, AvrSs50 | Srs5, Srs50 | Close to repetitive elements | (Clen et al., 2017; Salcedo et al., 2017) |
| *Blumeria graminis f. sp. hordei* | Barley | AvrK1, AvrA10, AvrA1, AvrA13, AvrA7, AvrA9, AvrA22 | M01, M01a, M01b, M01c, M01d, M01e, M01f, M01g, M01h, M01i | Close to repetitive elements | (Bourras et al., 2018; Saur et al., 2019) |
| *Blumeria graminis f. sp. tritici* | Wheat | AvrPn3a2/2, AvrPm2, AvrPm3b2/2, and AvrPm3d3 | Pn3a, Pn3b, Pm3a, Pm3b, Pm3c, Pm3d | Close to repetitive elements | (Bourras et al., 2019, 2018) |
| *Verticillium dahliae* | Tomato | Ave1 | – | Close to repetitive elements | (de Jonge et al., 2012) |
| *Avirulence proteins determining host species specificity* | *Magnaporthe oryzae* | Grass species | Pwl2, PWT3, PWT4 | Rws3 and Rws4 | – | (Inose et al., 2017; Sweigard et al., 1995; Takabayashi et al., 2002) |
| *Blumeria graminis f. sp. tritici* | Wheat | AvrPn3a2/2, AvrPm3d3 | Pm3b, Pm3c, Pm3d | Close to repetitive elements | (Bourras et al., 2019) |
| *Proteinaceous host-specific toxins determining host cultivar specificity* | *Stagonospora nodorum* | Wheat | SnTox1, SnTox2, SnTox3 | Snt1 and Snt2 | Repeat-rich regions | (Friesen et al., 2006; Liu et al., 2012, 2009) |
| *Pyrenophora tritici-repentis* | Wheat | T oxA and T oxB | T oxA and T oxB | Repeat-rich regions | (Friesen et al., 2006; Martinez et al., 2004) |
| *Secondary metabolites determining host cultivar specificity* | *Cochliobolus carbonum* | Corn | HC-toxin | Hm/hm | Repeat-rich region | (Walton, 2006) |
| *Cochliobolus heterostrophus* | Corn | T-toxin | T -toxin | Repeat-rich regions | (Inderbitzin et al., 2010; Yang et al., 1996) |
| *Secondary metabolites determining host species specificity* | *Verticillium dahliae* | Cotton, olive and okra | VdDf5 and VdDf6 | – | Lineage-specific genomic region | (Clen et al., 2018; Zhang et al., 2019) |
| *Alternaria alternata* | A number of crops | AM-toxin, AF-toxin, AK-toxin, ACT-toxin, ACR-toxin, AAL-toxin, AT-toxin | – | Accessory chromosome | (Akimitsu et al., 2014; Tsuge et al., 2013) |
...effector from the wheat tan spot pathogen (Wang et al., 2015). These two effectors, together with AvrPiz-t and ToxB, an protein family has no homology to proteins with known function. Furthermore, it shows homology with the nudix hydrolase superfamily and is the first protein containing a Pfam domain. AvrM14-A (Anderson et al., 2016), respectively. AvrL567-B is recognized most strongly by L5, weakly by L6 and not at all by L7 (Dodds et al., 2004). The gene encoding AvrL567-C co-segregates with the virulence phenotype, and this version is not recognized by L5, L6, or L7 (Dodds et al., 2004). AvrP4 and AvrP123 are cysteine-rich proteins, whereas AvrM does not contain cysteine residues at all. AvrL2-A and AvrM14-A are identified by map-based cloning (Anderson et al., 2016). AvrM14-A is not related to AvrM and is recognized by both the flax M1 and M4 resistance proteins (Anderson et al., 2016). AvrM14-A shows homology with the nudix hydrolase superfamily and is the first rust avirulence protein for which a biochemical function could be predicted from the protein sequence (Anderson et al., 2016). The AvrL2 protein family has no homology to proteins with known function.

The three-dimensional structures of AvrL567, AvrM and AvrP have been determined that infects flax (Linum usitatissimum) and other species of the genus Linum (Lawrence et al., 2007). To date, avirulence genes have been identified in six loci in M. lini, encoding AvrL567 (Dodds et al., 2004), AvrM (Catanzariti et al., 2006a), AvrP123 (Catanzariti et al., 2006b), AvrP4 (Catanzariti et al., 2006b), AvrL2-A (Anderson et al., 2016) and AvrM14-A (Anderson et al., 2016), respectively. AvrL567-A is recognized by resistance proteins L5, L6 and L7, whereas AvrL567-B is recognized most strongly by L5, weakly by L6 and not at all by L7 (Dodds et al., 2004). The gene encoding AvrL567-C co-segregates with the virulence phenotype, and this version is not recognized by L5, L6, or L7 (Dodds et al., 2004). AvrP4 and AvrP123 are cysteine-rich proteins, whereas AvrM does not contain cysteine residues at all. AvrL2-A and AvrM14-A are identified by map-based cloning (Anderson et al., 2016). AvrM14-A is not related to AvrM and is recognized by both the flax M1 and M4 resistance proteins (Anderson et al., 2016). AvrM14-A shows homology with the nudix hydrolase superfamily and is the first rust avirulence protein for which a biochemical function could be predicted from the protein sequence (Anderson et al., 2016). The AvrL2 protein family has no homology to proteins with known function.

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acid protein which interacts with Sr50 directly (Chen et al., 2017; Salcedo et al., 2017). The origin of isolates virulent on Sr25-containing plants is associated with the insertion of a miniature inverted transposable element (MITE) in AvrSr35 (Salcedo et al., 2017). A switch to virulence towards Sr50 was due to the exchange of a whole chromosome between two haploid nuclei, resulting in loss of the avirulence allele (Chen et al., 2017). In addition, the Pgt protein PGTAUSPE-10-1 causes cell death in a host line carrying resistance gene Sr22. Therefore, PGTAUSPE-10-1 might be the avirulence factor corresponding to Sr22 (Upadhyaya et al., 2014). Through mutagenic genomics approaches, AvrSr27 has also been identified, and identification of AvrSr5 is underway (Dodds et al., 2019).

2.8. Blumeria graminis

2.8.1. B. graminis f. sp. hordei (Bgh)

B. graminis f. sp. hordei (Bgh) causes powdery mildew on barley, and interacts with its host in a gene-for-gene manner (Zhang et al., 2005). So far, seven avirulence genes have been identified in Bgh, namely AvrK1, AvrA1, AvrA7, AvrA9, AvrA10 and AvrA22, corresponding to barley resistance genes mla1, mla13, mla7, mla9, mla10 and mla22, respectively (Lu et al., 2016; Ridout et al., 2006; Saur et al., 2019). AvrA7, AvrA9, AvrA10, and AvrA22 were identified in a recent study and these proteins interact directly with their respective plant R proteins (Saur et al., 2019). Virulence of isolates is predominately associated with non-synonymous SNPs and loss of expression of avirulence genes (Saur et al., 2019). Ava10 and Ava22 share homology and associated with non-synonymous SNPs and loss of expression of avirulence genes (Saur et al., 2019). AvrA10 and AvrA22 probably have evolved through two opposing selective pressures: sequence conservation and sequence diversification to maintain their virulence function and sequence diversification to escape recognition by Avr10 and Avr22, respectively (Saur et al., 2019). Avra7, Avra9, Avra10, and Avra22 were identified in a recent study and these proteins interact directly with their respective plant R proteins (Saur et al., 2019).

2.8.2. B. graminis f. sp. tritici (Bgt)

B. graminis f. sp. tritici (Bgt) is the causal agent of powdery mildew on wheat (Bourras et al., 2018). Four genes encoding avirulence factors have been identified in Bgt: AvrPm3\(^{2/2}/2\), AvrPm2, AvrPm3\(^{2/2}/2\) and AvrPm3\(^{2/2}/2\). The encoded products are recognized by R proteins Pm3a/3f, Pm2, Pm3b/3c and AvrPm3d, respectively (Bourras et al., 2019, 2015; Praz et al., 2017). The avirulence genes all encode small, secreted proteins with conserved cysteine residues, and they all are highly expressed in haustoria (Bourras et al., 2019, 2015; Praz et al., 2017). AvrPm2 belongs to a small gene family encoding structurally conserved RNAse-like effectors (Praz et al., 2017). Recognition of AvrPm3\(^{2/2}/2\), AvrPm3\(^{2/2}/2\) and AvrPm3\(^{2/2}/2\) by the respective resistance proteins is suppressed by the ribonuclease-like effector SvrPm3\(^{1/1}/1\) (Bourras et al., 2019). AvrPm3\(^{2/2}/2\) shows a high level of copy number variation in mildew isolates, while AvrPm3\(^{2/2}/2\) is present as a single copy only (Bourras et al., 2019). Although Pm3 alleles share more than 97% sequence identity on the protein level, AvrPm3\(^{2/2}/2\), AvrPm3\(^{2/2}/2\) and AvrPm3\(^{2/2}/2\) share low sequence identity (Bourras et al., 2019). In the fungal genome all these avirulence genes are surrounded by TEs (Bourras et al., 2019, 2015; Praz et al., 2017). In 185 isolates, 10 non-synonymous mutations were found across the AvrPm3\(^{2/2}/2\) gene, while 17 non-synonymous mutations were found for AvrPm3\(^{2/2}/2\) (Bourras et al., 2019). AVRPm3\(^{2/2}/2\) and AVRPm3\(^{2/2}/2\) homologs have also been found in rye and Dactylis medicws. The products of the homologs are recognized by the resistance proteins Pm3b, Pm3c and Pm3d in wheat, demonstrating that AvrPm3-Pm3 interactions also determine host species specificity in cereal mildews.

2.9. Verticillium dahliae

V. dahliae is an asexual soil-borne, xylem-invading plant pathogen that causes vascular wilt diseases in over 200 dicotyledonous plant species, such as tomato (Klosterman et al., 2009). So far, identification of only one avirulence protein, Ave1, has been published. Ave1 is a 134 aa secreted protein recognized by Ve1 in tomato, and expression of Ave1 is induced during host colonization (de Jonge et al., 2012). Intriguingly, no SNP was found in 85 Ave1 alleles from Verticillium strains isolated from various host plants and different geographical locations (de Jonge et al., 2012). The presence of numerous Ave1 orthologs in plants, absence of orthologs in fungi other than Fol, Colletotrichum higginsianum, and Cercospora betollia, and the association of Ave1 with a flexible genomic region containing various TEs suggest that Verticillium acquired Ave1 from plants through horizontal gene transfer (HGT) (de Jonge et al., 2012). In a recently study, Ave1 was shown to play a role in niche colonization by suppressing microbes with antagonistic activities (Snelders et al., 2020).

In another study of V. dahliae (personal communication with Jinling Li, unpublished data), a duplicated defoliation-specific gene, encoding a small secreted protein, was found. It was demonstrated that this effector gene is required for the defoliation of cotton and olive. Application of this heterologously produced protein to cotton seedlings also induced defoliation, indicating that the protein is directly responsible for the defoliation symptoms.

3. Virulence genes determining host cultivar specificity

3.1. Stagonospora nodorum

The ascomycete fungus S. nodorum (also known as Parastagonospora nodorum) is a major necrotrophic pathogen of wheat causing leaf and glume blight (Liu et al., 2009). So far, three S. nodorum genes encoding host selective toxins, notably SnToxA (Liu et al., 2012), SnTox1 (Liu et al., 2009) and SnTox3 (Liu et al., 2009) have been identified. The genes reside on three different chromosomes (Liu et al., 2012). The toxins induce cell death and necrosis as an outcome of their interaction with their cognate dominant susceptibility gene products (ToxA-Tsn1, Tox1-Snn1 and Tox3-Snn3). Below these host-specific interactions are summarized.

3.1.1. SnToxA-Snn1

Although the SnToxA-Sn1 interaction was the first to be characterized (Liu et al., 2004), the toxin gene was identified only in 2012 (Liu et al., 2012). The mature SnTox1 contains 100 amino acids including 16 cysteine residues, all predicted to be involved in disulfide bridges necessary for the activity/stability of the protein. The C-terminus shows similarity to the chitin-binding domain of Avr4 of C. fulvum (van Esse et al., 2007). Later, it was shown that SnTox1 binds chitin of the fungal cell wall, protecting the pathogen from chitinase degradation (Liu et al., 2016). SnTox1 is highly expressed at 3 days post infection (dpi), which correlates with the onset of necrotic lesion development. Diferent from other effector genes in S. nodorum, which are located in gene poor and repeat-rich regions, SnTox1 is located in a gene-rich region and no obvious repeats or AT-rich sequences were identified within the 300 kb region containing SnTox1 (Liu et al., 2012). Among 159 Sn isolates from around the globe, 11 Tox1 isoforms were found, suggesting diversifying selection on SnTox1 (Liu et al., 2012).

3.1.2. SnToxA-Tsn1 and PtrToxA-Tsn1

SnToxA is highly similar to PtoxA identified in P. tritici-repentis (Ciuftetti et al., 1997), having only two amino acid differences, and both are recognized by the wheat protein Tsn1 (Faris et al., 2010). Mature SnToxA is a 13.2 kDa protein containing two cysteine residues as well as an RGD-containing vitronectin-like motif that is present in a solvent-exposed loop in the active protein (van Esse et al., 2007). The regions upstream and downstream of SnToxA contain repetitive, AT-rich regions, but these AT-rich regions have not been found in the corresponding flanking regions of P. tritici-repentis ToxA.

By sequencing 95 S. nodorum ToxA and 54 P. tritici-repentis ToxA
3.2. Pyrenophora tritici-repentis (P. tritici-repentis)
resulting in four amino acid polymorphisms (Liu et al., 2009). The gene has likely undergone HGT from S. nodorum to P. tritici-repentis based on the following observations: (1) the presence of an almost identical 11 kb region containing SnToxA/PtrToxA in both species; (2) the presence of high sequence diversity of SnToxA in S. nodorum and monomorphism in Ptr; (3) absence of PtrToxA in related species; (4) the recent emergence of tan spot (Friesen et al., 2006; Liu et al., 2006).

3.1.3. SnTox3-Snn3
SnTox3 was identified by partial purification and sequencing of the protein (Liu et al., 2009). SnTox3 encodes a 230 amino acid pre-pro-protein consisting of a 20 amino acid signal sequence and a predicted pro-domain of approximately 30 amino acids, resulting in a mature protein of ~18 kDa (Liu et al., 2009). SnTox3 contains six cysteine residues, each being predicted to be involved in the formation of a disulfide bridge critical to the structure and function of the protein (Liu et al., 2009). Like SnTox1, SnTox3 is highly expressed at 3 dpi when lesions start to develop. SnTox3 interacts with a wheat pathogenicity-related-1 protein (Breen et al., 2016). SnTox3 is also flanked by AT-rich sequences, containing long terminal repeat retrotransposons. By sequencing SnTox3 from 245 isolates, eleven haplotypes were identified resulting in four amino acid polymorphisms (Liu et al., 2009).

3.2. Pyrenophora tritici-repentis (P. tritici-repentis)
The ascomycete Pyrenophora tritici-repentis (Ptr) causes tan spot and chlorosis on wheat. Two Ptr host-specific genes encoding host-specific toxins have been identified, including Ptr-ToxA (Giuffetti et al., 1997) and Ptr-ToxB (Strelkov et al., 1999). Ptr-ToxA and Ptr-ToxB interact specifically with the products of the host susceptibility genes Tsn1 and Tsc2, respectively. It is likely that Ptr-ToxA originated from S. nodorum (see above) (Friesen et al., 2006). Ptr-ToxA is a single domain protein having a β-sandwich fold with two antiparallel β-sheets composed of four strands each enclosing the hydrophobic core (Sarma et al., 2005). Unlike Ptr-ToxA, which is present as a single copy in the genome, Ptr-ToxB is present in multiple copies in the genome of some races of Ptr, and the number of copies is proportional to virulence (Martinez et al., 2004). Ptr-ToxB encodes a 64 amino acid host-selective toxin (Martinez et al., 2001), contains four cysteine residues involved in the formation of two disulfide bridges (Nyarko et al., 2014) and is a heat-stable protein (Strelkov et al., 1999). All Ptr-ToxB loci are associated with retrotrotransposons (Martinez et al., 2004). Ptr-toxb is a related single copy gene from a non-pathogenic strain and shares 86% identity with Ptr-ToxB (Martinez et al., 2004). Both Ptr-ToxB and Ptr-toxb adopt a β-sandwich fold stabilized by two disulfide bonds, but differ in the dynamics of one sandwich half. The absence of toxic activity of Ptr-toxb is due to production of HC-toxin, a cyclic tetrahydrofuran (THF) ring (Martinez et al., 2004). Both Ptr-ToxB and Ptr-toxb have a single locus (Panaccione et al., 1999). It contains HST1, encoding the 570 kDa non-ribosomal peptide synthetase key enzyme. TOX2 is duplicated in toxin-producing isolates of the fungus, but is completely absent from the genomes of toxin-non-producing isolates (Panaccione et al., 1999). Disruption of all copies of HST1 resulted in abolished HC-toxin production and loss of host-selective pathogenicity. Other genes involved in the processing of biosynthetic intermediates and possibly secretion include TOXA (major facilitator superfamily transporter) (Pitkin et al., 1996), TOXC (fatty acid synthase subunit) (Ahn and Walton, 1997) and TOXF (amino acid transaminase) (Cheng et al., 1999). Except for one copy of the pathway-specific regulator gene TOXE, all other TOX2 genes are located in an approximately 600 kb repeat-rich region (Ahn and Walton, 1996; Condon et al., 2013).

4. Cochliobolus heterostrophus
Cochliobolus heterostrophus (C. heterostrophus) causes Southern Corn Leaf Blight. There are two known races, race T and race O, with race T producing a HST called T-toxin. Race T is highly virulent on corn carrying Texas male sterile cytoplasm, while Race O only shows mild virulence. So far, nine genes have been shown to be required for T-toxin production (Baker et al., 2006; Inderbitzin et al., 2010; Rose et al., 2002; Yang et al., 1996), and these genes are located at two unlinked loci, designated as ToxA1 and ToxB1. Appropriately, these genes are absent in Race O. The nine known Tox1 genes encode two polyketide synthases (PKS), a decarboxylase, five dehydrogenases and an unknown protein (Inderbitzin et al., 2010). The genes do not reside in a single cluster, but reside alone or in small groups in four distinct AT-rich regions (Inderbitzin et al., 2010). Together, these Tox1 regions comprise less than 5% of the 1.2 Mb of race T-specific DNA (Inderbitzin et al., 2010).

4.3. Verticillium dahliae
As mentioned above, V. dahliae is a vascular wilt pathogen which can infect nearly 200 plant species. However, it causes defoliation in a few hosts only, including cotton, olive and okra (Milgroom et al., 2016). Recently, by using comparative genomics, seven genes associated with the defoliating pathotype, VdDf1-VdDf7, were discovered in a lineage-specific genomic region (G-LSR2) of V. dahliae Vd991 (Chen et al., 2018; Zhang et al., 2019). VdDfs and VdDf6 are critical for the defoliation phenotype (Zhang et al., 2019). The VdDfs are involved in the production of NAE 12-0, which cause defoliation either by altering abscisic acid sensitivity, hormone disruption or sensitivity to the pathogen (Zhang et al., 2019). Phylogenetic analysis of the region comprising all seven protein-coding genes suggests that G-LSR2 has been acquired from F. oxysporum f. sp. vasinfectum through horizontal transfer (Chen et al., 2018).

4.4. Alternaria alternata
Alternaria alternata (A. alternata) is a ubiquitous, mostly saprophytic fungus present in dead plant material but is also known as a weak pathogen causing opportunistic diseases in a number of crops (Akimitsu et al., 2014). HSTs in A. alternata have been extensively reviewed before (Akimitsu et al., 2014; Meena et al., 2017). So far, there are seven known diseases caused by A. alternata in which HSTs are responsible for pathogenesis. Accordingly, A. alternata is classified into seven different pathotypes, each producing a distinct host-specific toxin: AM-toxin (apple pathotype), AF-toxin (strawberry pathotype), AK-toxin (Japanese pear pathotype), ACT-toxin (tangerine pathotype), ACR-toxin (rough lemon pathotype), AAL-toxin (tomato pathotype), or AT-toxin (tobacco pathotype). Interestingly, HST genes in A. alternata are located on conditionally dispensable chromosomes (CDCs) as gene clusters (Hu et al., 2012). Chemical structures of HSTs from six pathotypes have been determined, excluding that of AT-toxin of the tobacco pathotype. AAL-toxin is a structural analog of sphingolipid precursors, and thereby efficiently inhibits eukaryotic ceramide synthases. Appropriately,
resistance in tomato is conferred by the ceramide synthase Asc-1 encoded within the *Alternaria* stem canker (Asc) locus (Abbas et al., 1994; Brandwagt et al., 2000; Spassieva et al., 2002).

5. Horizontal gene and chromosome transfer and host specificity

HGT or horizontal chromosome transfer (HCT) has likely occurred in several plant pathogenic fungi, including *Colletotrichum gloeosporioides* (He et al., 1998), *S. nodorum/P. tritic-repentis* (Friesen et al., 2006), *F. oxysporum* (Ma et al., 2010; van Dam et al., 2017; Vlaardingerbroek et al., 2016) and *A. alternata* (Akimitsu et al., 2014).

The outbreak of tan spot disease on wheat in 1941 was caused by *P. tritic-repentis*. The high virulence of *P. tritic-repentis* is due to the presence of a HST, ToxA, and, as mentioned above, it is likely that the ToxA gene was transferred from *S. nodorum* to *P. tritic-repentis* in a recent event (Friesen et al., 2006).

In *F. oxysporum*, HCT of pathogenicity chromosomes has been shown in three *formae specialiae*: Fol (Ma et al., 2010; Vlaardingerbroek et al., 2016), *F. oxysporum* f. sp. *radicis-cucumerinum* (Forc) (van Dam et al., 2017) and *F. oxysporum* f. sp. *melonis* (Fom) (Li et al., 2020). In all three cases, following transfer of an accessory chromosome containing effector genes from a pathogenic strain to non-pathogenic *F. oxysporum*, the recipient strain becomes pathogenic to the host species of the chromosome donor strain.

As mentioned above, in *A. alternata*, all HST-encoding genes are located on CDCs (Mehrabi et al., 2011). It has been suggested that these CDCs are transferrable between different pathotypes in *A. alternata* (Akagi et al., 2009). Indeed, HCT between different pathotypes has been demonstrated through protoplast fusion experiments. For example, by fusion of a tomato pathotype with a strawberry pathotype, the resulting strain was found to be pathogenic on both tomato and strawberry (Akagi et al., 2009).

Often, virulence or host-determining genes are clustered, so HGT or HCT can cause previously non-pathogenic microbes to become pathogenic, or pathogens to expand or change host range. In *A. alternata* and *F. oxysporum*, genes determining host range – a secondary metabolite gene cluster or virulence genes together with transcription factors – are located on a single chromosome, therefore single chromosome transfer is sufficient to expand host range. This may be an important mechanism for asexual fungi to generate genetic variation and adapt to a changing environment.

6. Potential role of genome localization of host-specificity genes in adaptation

From the findings discussed above, host-specificity genes appear to be predominately located in AT-rich isochores (such as avirulence genes in *L. maculans* and *AvrPm3a*/*b* and *AvrPm3c* in *B. graminis* f. sp. *tritici*), and/or TE-rich lineage-specific chromosomes (such as AVR1, AVR2, and AVR3 in Fol). These repeat-rich genomic compartments are believed to evolve more rapidly than other parts of the genome. Based on this, the concept of a ‘two-speed genome’ has been proposed (Croll and McDonald, 2012; Raffaele and Kamoun, 2012). The gene-poor, repeat-rich genomic compartment could serve as a ‘cradle’ for adaptive evolution (Croll and McDonald, 2012). Evidence for a higher diversification rate has been found for almost all the host-specificity genes listed above. For example, in *L. maculans*, avirulence genes are located in AT-rich isochores and these genes have an exceptionally high mutation rate due to RIP (Rouxel and Balesdent, 2017).

Modern crop management practices have accelerated the arms race between pathogens and plants (Möller and Stukenbrock, 2017). Cultivars with new resistance genes may be introduced in each growing season, which poses great pressure on pathogens to rapidly adapt to the new cultivar. Since avirulence factors that are recognized by resistance proteins are generally located in repeat-rich genomic regions, a high frequency of point mutations, deletions, duplications, silencing or rearrangement of avirulence genes can result in rapid emergence of strains that evade recognition by resistance proteins. If loss of or changes in avirulence genes have no or little fitness cost, these strains will quickly become dominant in the pathogen population. The compartmentalization of a genome thus allows pathogens to harbor fast-evolving genes without affecting the stability of core genes.

7. Conclusions and perspectives

Both proteins and secondary metabolites can determine host specificity in plant pathogenic fungi. With an increased rate of publication of pathogen genome sequences, the rate of discovery of genes determining host cultivar-specificity will likewise increase, by comparing strains of different races within a species or *forma specialis*. Identification of host species-specificity factors may require more effort, depending on the fungal species involved. Regarding the latter, it would be interesting to test the hypothesis of Schulze-Lefert and Panstruga, who proposed that PTI plays a major role in non-host resistance of evolutionary divergent non-host plant species, while ETI would be more dominant as a non-host resistance mechanism in more closely related plant species (Schulze-Lefert and Panstruga, 2011). To test whether ETI indeed plays a dominant role in non-host resistance of closely related plant species, we propose to identify effectors eliciting an ETI response in non-host plant species closely related to host species in fungal pathogens such as *M. oryzae* or *F. oxysporum*.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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