Perspectives on the utilization of resistance mechanisms from host and nonhost plants for durable protection of *Brassica* crops against Alternaria blight

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**ABSTRACT**

**Background.** *Alternaria brassicae*, the causal organism of Alternaria blight, is a necrotroph infecting crops of the *Brassicaceae* family at all growth stages. To circumvent this problem, several disease management strategies are being used in the field, and disease-resistant varieties have also been developed. However, no strategy has proven completely successful, owing to the high variability in virulence among *A. brassicae* isolates, which causes a diverse spectrum of symptoms. Nonhost resistance (NHR) is a robust and broad-spectrum defense mechanism available in plants, and the exploitation of gene pools from plant species that are nonhost to *A. brassicae* could serve as novel sources of resistance.

**Methodology.** We searched the literature using key words relevant to this study in various search engines, such as PubMed, Web of Science, and Google Scholar, as well as certain journal websites. The literature was retrieved, sorted, and mined to extract data pertinent to the present review.

**Results.** In this review, we have comprehensively covered the recent progress made in developing Alternaria blight resistance in *Brassica* crops by exploiting host germplasm. We also enumerate the potential NHR sources available for *A. brassicae* and the NHR layers possibly operating against this pathogen. In addition, we propose different strategies for identifying NHR-related genes from nonhost plants and testing their relevance in imparting broad-spectrum resistance when transferred to host plants.

**Conclusion.** This review will help broaden the current knowledge base pertaining to the resistance sources available in host germplasm, the exploitation of NHR mechanisms, and their applications in protecting *Brassica* crops from Alternaria blight. The insights might also be applicable to a wider repertoire of plant pathogens.

**Subjects** Agricultural Science, Molecular Biology, Mycology, Plant Science

**Keywords** Brassica, Alternaria Brassicae, Broad-spectrum defense, Blight-disease, Necrotrophic fungus, Nonhost resistance
INTRODUCTION

Despite the considerable increase in the production and productivity of Brassica crops (B. juncea, B. rapa, B. napus), there remains a large discrepancy between yield potential and crop productivity in farmers’ fields owing to the plethora of diseases affecting these crops. Among these diseases, Alternaria blight, caused by the necrotrophic fungus Alternaria brassicae, is one of the most important diseases of oilseed brassicas throughout the world (Conn, Tewari & Awasthi, 1990; Saharan, Mehta & Meena, 2016). Two other species—A. brassicicola and A. raphani—are also involved in disease manifestation across the world. However, A. brassicae is highly infectious and more damaging to Brassica crops (Kadian & Saharan, 1983; Meena et al., 2010). A. brassicae can affect the host plant at all stages of growth, including seeds. Infected seeds and diseased plant debris are the sources of primary infection. However, under favorable environmental conditions, the pathogen spreads through airborne and soil-borne spores, resulting in secondary infection. A. brassicae has no known sexual stage and has been found to survive as mycelium or spores on decaying plant debris or as latent infection in seeds (Thomma, 2003). Under temperate conditions, the pathogen survives in the form of mycelia or conidia on the previous year’s crop debris and as chlamydospores or microsclerotia at cooler temperatures (Tsuneda & Skoropad, 1977; Humpherson-Jones & Maude 1982; Humpherson-Jones & Phelps, 1989). It also resides in alternative hosts such as susceptible weeds or perennial crops during the off season (Chupp & Sherf, 1960; Maude & Humpherson-Jones, 1980). The spores landing on the host surface as conidia adhere to the surface and penetrate the host mainly through the stomata after forming a germ tube, which then grows into the host epidermal cells through the formation of an appressorium, often triggered by host signaling mechanisms (Yang et al., 2005; Cho et al., 2006). The infection results in the appearance of severe disease symptoms, including numerous black spots covering the pods, leading to losses in seed yield of up to 60% (Kolte, 1985; Kolte, 2002; Meena et al., 2010). The severity of the disease differs among seasons and regions, as well as among crops within a region (Meena et al., 2012), which can be attributed to the variability among isolates of A. brassicae species. Different strategies have been adopted to control the disease, such as crop rotation practices and fungicide application; however, the devastating effect of this pathogen on yield loss is not effectively overcome. Among the different strategies, the most viable approach has been the utilization of resistant sources from host germplasm (Meena et al., 2010). The transfer of resistance from tolerant wild species of the Brassicaceae family to cultivated Brassica crops has been repeatedly attempted but has yielded limited success owing to variability in virulence among A. brassicae isolates.

Another potential strategy parallel to this could be the transfer of genes from nonhost plants conferring resistance to all the isolates of A. brassicae. Nonhost resistance (NHR) is the most durable form of plant immunity effective against all genetic variants of a pathogen (Heath, 2000; Mysore & Ryu, 2004; Senthil-Kumar & Mysore, 2013; Lee et al., 2017). NHR is multilayered and can be divided into two major types: the pre-invasion layer and the post-invasion layer (Nurnberger & Lipka, 2005; Lee et al., 2017; Fonseca & Mysore, 2018). The pre-invasion layer has both pre-formed and inducible defense mechanisms.
Pre-formed defense involves the physical and chemical barriers that prevent fungal entry inside the plant (Lee et al., 2017; Fonseca & Mysore, 2018). In addition, the plant recognizes pathogen-associated molecular patterns (PAMPs) and elicitor molecules from fungal pathogens and induces defense responses against them. Meanwhile, the post-invasion layer mainly comprises inducible defense responses from the plant. In the post-invasion layer, the defense response culminates in a hypersensitive response (HR) and reactive oxygen species (ROS)-mediated cell death. These defense responses in the post-invasion layer can further restrict fungal growth inside the plant (Lee et al., 2017; Fonseca & Mysore, 2018). These NHR layers together offer durable resistance and can be utilized for developing disease-resistant crops. This calls for the identification of the genes involved in NHR and the elucidation of the NHR mechanisms against A. brassicae. Eventually, NHR-related genes can be transferred to host plants to confer durable resistance to A. brassicae. In this review, we provide insights on the resistance mechanisms operating in Alternaria blight-tolerant Brassica species and the underlying NHR mechanisms. We also propose the potential strategies for exploiting NHR mechanisms by using genomic tools for future protection of Brassica crops.

**Survey methodology**

Several search engines, including PubMed, Web of Science advanced search, and Google Scholar, as well as specific journal websites, were used, and the search was performed based on key words specifically chosen for the topic. Literature was retrieved and sorted based on the relevance of the topic. The literature was then mined to extract data, and relevant articles published from 1960 to 2018 were used to support and elaborate the hypothesis of this review. Various citations from these articles were back-referred to obtain further detailed information. Together, the compiled information was processed by the authors to prepare the manuscript. Perspectives on the relevant concepts were incorporated based on the authors’ expertise in this field of research.

**Symptoms and epidemiology**

Alternaria blight disease symptoms occur in the leaves, petiole, inflorescence, stem, pods, and seeds (Verma & Saharan, 1994). The initial symptoms of blight appear on the older leaves of the plant, as discrete pinpoint spots, which later enlarge and become surrounded by a distinct yellow halo with concentric rings (Kumar et al., 2014). These spots ultimately coalesce, forming large patches of chlorotic and necrotic foliar blight lesions, resulting in defoliation. The pathogen resides in the center of the lesions, which is surrounded by yellow halos, the zone created by the diffusion of fungal metabolites (Tewari, 1983; Agarwal et al., 1997). Symptoms produced by all three species of Alternaria are similar. However, the spots formed by A. brassicae are greyish in color (Kumar et al., 2014). On the stems, the spots are elongated, while they appear round and blackish on the pods. Infection on the leaves and pods reduces photosynthetic potential, which not only results in reduced seed development and weight but also in reduced oil content and quality and poor germination efficiency (Kolte, 1996; Meena et al., 2010). In severely infected plants, the entire pod gets covered with numerous black spots, ultimately causing significant yield loss due to premature shattering.
In field conditions, disease development and spread are generally favored under temperatures ranging from 12 °C to 27 °C and a relative humidity of 70%, while on pods, infection occurs at a daily temperature range of 20–30 °C, together with more than 9 h of sunshine and leaf wetness (Awasthi & Kolte, 1994). Frequent rains are favorable for disease initiation and spread on the leaves of Brassica, particularly during the rosette to flowering stages. Furthermore, the frequency of infection is the highest during the flowering and pod stages (Prasad, Saxena & Chandra, 2003). According to a report by Awasthi & Kolte (1994), the susceptibility of the plants increases with age 30 days after sowing. Plants less than 30 days old do not show any symptoms, but they are highly susceptible 60–90 days after sowing.

**Resistance mechanisms operating in wild and tolerant species of Brassica**

Various studies have been carried out to analyze the effect of Alternaria blight infection on host plant physiology and metabolism, but reports on host–pathogen interactions at the genetic and molecular level are still limited. A recent study has indicated that Arabidopsis can be used as an efficient model system for deciphering the genetic and molecular mechanisms involved during interaction with A. brassicae (Mandal, Rajarammohan & Kaur, 2018).

Studies on the mechanisms of resistance to Alternaria blight have implicated polygenes (Tripathi et al., 1980; Medhi, 1985; Zhang, Xu & Takahata, 1996; Krishnia, Saharan & Singh, 2000), while some studies have attributed it to dominant nuclear genes (Tripathi et al., 1978; Subudhi & Raut, 1994; Zhang et al., 1997). Thus far, different sources of resistance to A. brassicae have been identified from host germplasm, including wild and tolerant species of Brassica. The resistance in the host germplasm has different components, which mainly include structural components such as epicuticular waxes, as well as biochemical components like phenols and phytoalexins. Epicuticular waxes form a direct physical barrier in the plant, providing resistance to Alternaria (Conn, 1986; Conn & Tewari, 1989). High deposits of epicuticular wax, forming a protective hydrophobic coating on the leaf surface, reduce the adherence of inoculum, conidia germination, and germ tube formation in the plant (Saharan, 1992). Some Brassicaceae members, including B. napus, B. carinata, and Sinapis alba, have been reported to have higher epicuticular wax compared to B. rapa and B. juncea, and therefore, the former are less susceptible to Alternaria blight infection (Conn, Tewari & Hadziyev, 1984; Tewari, 1986). High quantities of epicuticular wax have been observed in the progeny of interspecific crosses between B. napus and B. juncea (Singh, Singh & Srivastava, 1999).

Alternaria blight-resistant varieties accumulate high amounts of biochemical compounds such as phenols. After infection, blight-tolerant species of Brassica such as B. carinata and B. napus have been observed to accumulate a higher amount of total phenols compared to the susceptible species B. juncea and B. rapa (Gupta et al., 1990; Gupta, Gupta & Kaushik, 1995; Gupta & Kaushik, 2002). Meanwhile, the levels of soluble and reducing sugars and soluble nitrogen have been found to be lower in resistant species. In another study, resistance to the disease was reported to be associated with increased levels of
antioxidant enzymes of the phenolic pathway, such as polyphenol oxidase, peroxidase, and catalase (Singh et al., 2009). Furthermore, phytoalexin accumulation in response to pathogen infection and its role in disease resistance have been well studied in Brassica. Phytoalexins are low-molecular-weight antimicrobial compounds produced by plants after pathogen infection. Cultivars of both resistant (B. napus, Camelina sativa, Eruca sativa) and susceptible (B. rapa) species show the elicitation of phytoalexins (Tewari, Conn & Dahiya, 1987; Tewari, Conn & Dahiya, 1988; Conn, Tewari & Dahiya, 1988; Conn et al., 1991). The highly resistant C. sativa produces a large number of phytoalexins that are involved in regulating resistance to A. brassicae (Tewari, 1991). Rapid accumulation of phytoalexins in C. sativa after pathogen infection has been found to inhibit fungal growth on the leaf surface (Jejelowo, Conn & Tewari, 1991).

The Alternaria pathogen has also been found to secrete both host-specific and nonhost-specific toxic metabolites, enabling a wide range of infection symptoms (Nishimura & Kohmoto, 1983; Kohmoto, Otani & Tsuge, 1995; Agrios, 2005). These toxins have been shown to alter the permeability and functioning of the cell membrane and organelles, thereby inhibiting various physiological processes of the host plant (Mathur & Chand, 1991). Destruxin B, a host-specific Alternaria phytotoxin acts as a pathogenicity factor responsible for its aggressiveness and for the susceptibility of the host plant. The Alternaria-tolerant species S. alba detoxifies destruxin B, and this is followed by simultaneous phytoalexin formation and elicitation. Together, these processes constitute the resistance mechanism of S. alba against A. brassicae (Pedras & Smith, 1997; Pedras et al., 2001). Some compounds related to camalexin and 6-methoxycamalex have also been found to confer toxicity to A. brassicae (Dzurilla et al., 1998).

**Exploiting the resistance mechanisms from host germplasm for developing blight-resistant brassicas**

Several attempts have been made to exploit resistance sources from the wild relatives of Brassica and transfer them into cultivated Brassica (Aneja & Agnihotri, 2013). Efforts have been made in screening blight-resistant cultivars; however, high levels of resistance to A. brassicae have not been observed among cultivated Brassica species. The species show varying degrees of resistance responses against Alternaria blight, and accordingly, B. rapa and B. juncea are classified as being more susceptible to the disease than B. napus and B. carinata (Skoropad & Tewari, 1977). Of late, studies aimed at identifying blight-tolerant oilseed Brassica germplasm have been increasing. Table S1 enlists the sources of resistance, including the related wild species or weedy varieties of Brassica against Alternaria blight. These related wild species have been found to harbor genes conferring resistance to Alternaria blight that can be used for resistance breeding (Ghose et al., 2008; Chatterjee, Mazumder & Basu, 2013).

In the past few decades, various conventional and modern approaches have been used to develop blight-resistant cultivars by exploiting resistance sources from host germplasm. Conventional approaches involve the screening of germplasm for blight-resistant plant materials, followed by the transfer of resistance traits from tolerant species into susceptible cultivars by hybridization combined with multiple crosses. Inter- and intraspecific crosses...
of B. juncea and B. carinata were established to understand the inheritance pattern of resistance to Alternaria blight (Krishnia, Saharan & Singh, 2000), which showed that dominant and additive genes govern the inheritance of resistance to Alternaria blight. Thus, a successful breeding method should utilize both the gene effects. However, there is a limitation in such breeding approaches owing to the low availability of suitable resistance sources/genes within the available germplasm of cultivated Brassica and insurmountable self-incompatibility barriers in combining the resistance traits from distantly related wild Brassica species. Modern biotechnological interventions involving tissue culture and genetic transformation approaches have also been established for the development of improved varieties. In vitro ovary culture and ovule culture have been attempted for transferring resistance traits from S. alba cv. Carine to B. napus cv. Brutor (Chevre et al., 1994) and Erucastrum cardaminoides to B. oleracea var. alboglabra (Mohanty et al., 2009). Intergeneric hybrids of B. campestris and B. spinescens have also been developed (Agnihotri et al., 1991). Moreover, the somatic hybridization technique has been applied for transferring the resistance traits from blight-tolerant species to susceptible ones, for example, S. alba to B. napus (Primard et al., 1988), B. carinata to B. juncea (Sharma & Singh, 1992), and B. nigra and B. oleracea (Jourdan & Salazar, 1993). Attempts have also been made for the introduction of somaclonal variations for the incorporation of disease resistance/tolerance to Alternaria blight through mutagenesis using gamma rays, ethyl methanesulfonate, and ethyl nitrosourea (Verma & Rai, 1980; Agnihotri et al., 2009). In addition, the defense response might be activated by treatment with biotic and abiotic agents for combating A. brassicae infection in oilseed brassicas. Vishwanath et al. (1999) showed that resistance in a susceptible cultivar of B. juncea against the extremely virulent A. brassicae isolate A and reasonably virulent isolate C, both from B. carinata, was induced after inoculating the plants with the avirulent A. brassicae isolate D. As another example, the application of β-aminobutyric acid on the leaves of B. carinata was found to induce resistance to A. brassicae (Chavan, Bhargava & Kamble, 2013). In future, the activation of the defense response without gene manipulation may act as a better substitute for the conventional use of fungicides. Unfortunately, till date, none of the conventional or modern biotechnological methods have been found to be feasible or stable in developing blight-resistant Brassica species. Under such circumstances, it is pragmatic to use novel sources of resistance from plant species that are not hosts to the pathogen. The transfer of genes from nonhost plants conferring resistance to the pathogen is a promising strategy in parallel to the current efforts towards the development of oilseed Brassica varieties resistant to Alternaria blight.

Exploiting nonhost disease resistance mechanisms

NHR is expressed by all the members of a plant species to a particular pathogen (Nurnberger & Lipka, 2005). Exploiting NHR has been suggested as an option for developing broad-spectrum disease resistance in plants (Ellis, 2006; Lee et al., 2017). In recent years, considerable progress has been made in understanding the interaction of several host species and A. brassicae, but there are few and indirect reports on studies examining the NHR mechanisms involved during the interaction of A. brassicae with nonhost plants. Understanding the mechanisms and identifying the genes involved in NHR to
A. brassicae may provide an opportunity to engineer transgenic Brassica plants that provide durable resistance to A. brassicae. For example, in recent years, some success in genetic transformation involving the transfer of resistance genes from nonhost plants into oilseed Brassicas has been reported. Mondal et al. (2003) found that the overexpression of tobacco chitinase conferred resistance to Alternaria leaf spot in transgenic Brassica, as demonstrated by the inhibitory effect of the transgenics on the hyphal growth of A. brassicae. They further reported the development of transgenic B. juncea (cv. RLM 198) constitutively expressing a class I basic glucanase of tomato, which arrested the hyphal growth of A. brassicae by 15–54% (Mondal et al., 2007). Moreover, the introduction of the chitin-binding lectin hevein from rubber plants into B. juncea led to resistance to A. brassicae (Kanrar et al., 2002). Such studies highlight the necessity to dissect the mechanisms and reveal the genes involved in NHR for the development of resistant cultivars.

For this, the first step is to identify potential nonhost plants that can reveal robust and exploitable mechanisms for suitable transfer to a host plant for imparting broad-spectrum resistance. Besides, the nonhost plant should be distinct from the non-adapted plant, where, for example, the fungal spores fail to adhere, deposit, and remain on the leaf surface for germination (Niks & Rubiales, 2002). Here, we discuss some of the nonhost plants for A. brassicae based on the available literature and the intercropping systems followed in different parts of world. In many intercropping systems, species of the Brassicaceae family are cultivated with certain other plant species in the same field for many decades (Singh, Kumar & Singh, 2010; Shekhawat et al., 2012; Ao & Saud, 2016; Ebrahimi et al., 2016; Jeromela et al., 2017). In the same field, A. brassicae infects Brassica plants but cannot infect the other plant species (http://www.plantwise.org/KnowledgeBank). The Plantwise Knowledge Bank database (http://www.plantwise.org/KnowledgeBank) lists the plant species infected by A. brassicae in different parts of world. Through this database, we ensured that the identified nonhost plants have not been reported to be infected by A. brassicae. Hence, those species can act as potential nonhost plants against this pathogen. Curiously, in the course of evolution, the pathogen gets enough opportunities to infect these nonhost plants, but they are unable to overcome the NHR. A few examples of such nonhost plants, which are intercropped with host plants such as mustard (B. juncea), are chickpea (Cicer arietinum and C. kabulium), lentil (Lens culinaris), cowpea (Vigna unguiculata), barley (Hordeum vulgare), sugarcane (Saccharum officinarum), and potato (Solanum tuberosum) (Table 1). It would be interesting to determine the factors that dictate the failure of A. brassicae to infect these nonhost plants. The NHR of these plants to A. brassicae may involve multiple layers of defense that could be distinct for each plant species.

### Possible NHR mechanisms in nonhost plants to A. brassicae

NHR to fungi is highly conserved among different plant species and seems to involve multiple layers of resistance (Heath, 2000). NHR mainly comprises pre-invasion and post-invasion defenses (Lee et al., 2017; Fonseca & Mysore, 2018). Based on the current understanding of the NHR mechanisms in other related nonhost plant–pathosystems such as Arabidopsis and powdery mildew fungus, barley and powdery mildew fungus, and Medicago and the rust fungal pathogen (Collins et al., 2003; Lipka et al., 2005; Jafary et al.,
### Table 1  List of the potential nonhost plants towards *A. brassicae*.

| Sl. No. | Name of the nonhost plant | Name of the host plant intercropped with nonhost plant\(^a\) | Criteria for categorization of nonhost plant | Information source |
|---------|---------------------------|---------------------------------------------------------------|-----------------------------------------------|--------------------|
| 1.      | Chickpea (*Cicer arietinum* and *C. kabalium*) | *B. juncea, B. napus B. rapa* | Intercropping system | *Gangasaran & Giri* (1985), *Tiwari et al.* (1992), *Ahlawat & Sharma* (2002), *Ahlawat, Gangaiyah & Ompal* (2005), *Singh, Kumar & Singh* (2010), *Shekhawat et al.* (2012), *Jeromela et al.* (2017) |
| 2.      | Lentils (*Lens culinaris*) | *B. juncea B. rapa* | Intercropping system | *Gangasaran & Giri* (1985), *Tiwari et al.* (1992), *Shyam, David & Philip* (2007), *Shekhawat et al.* (2012), *Ao & Saud* (2016), *Jeromela et al.* (2017) |
| 3.      | Barley (*Hordeum vulgare*) | *B. juncea B. rapa* | Intercropping system | *Gangasaran & Giri* (1985), *Singh, Kumar & Singh* (2010), *Shekhawat et al.* (2012) |
| 4.      | Wheat (*Triticum aestivum*) | *B. juncea B. napus B. rapa* | Intercropping system; Literature | *McRoberts & Lennard* (1996), *Ahlawat & Sharma* (2002), *Singh, Kumar & Singh* (2010), *Shekhawat et al.* (2012), *Ebrahimii et al.* (2016) |
| 5.      | Tomato (*Solanum Lycopersicum*) | – | Literature | *McRoberts & Lennard* (1996) |
| 6.      | Potato (*Solanum tuberosum*) | *B. juncea B. napus* | Intercropping system | *Ahlawat & Sharma* (2002), *Singh, Kumar & Singh* (2010), *Shekhawat et al.* (2012) |
| 7.      | Sugarcane (*Saccharum officinarum*) | *B. juncea, B. rapa, B. napus, B. campestris* | Intercropping system | *Singh, Kothari & Tripathi* (1986), *Toor et al.* (2000), *Suman et al.* (2006), *Ahlawat & Sharma* (2002), *Singh, Kumar & Singh* (2010) |

**Notes.**

\(^a\)Based on the information available on the plant wise knowledge bank database. ([http://www.plantwise.org/KnowledgeBank](http://www.plantwise.org/KnowledgeBank)) it was ensured that the above listed nonhost plants are not reported to be infected by *A. brassicae*. The intercropping system was considered as a criteria for the selection of the nonhost plant, when both the host and nonhost plants are cultivated together for many decades and the pathogen get enough chance to infect the nonhost plant. Other nonhost plants are selected from the literature studies based on experimental evidence that suggest the plant as a nonhost for *A. brassicae*.  

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...we have proposed the possible NHR mechanisms in a nonhost plant against *A. brassicae* (Fig. 1). The NHR mechanisms operating at the pre-invasion level mainly involve both pre-formed and inducible defense responses. The germination of *A. brassicae* spores requires host stimuli, high humidity, and surface water to increase wetness on the leaf surface (Verma & Saharan, 1994). Pre-formed defenses may involve structural features such as the abundance of leaf trichomes, which reduces leaf wettability, as well as the lack of stimuli from the nonhost plant, which eventually causes irregular spore germination or a complete lack thereof (Niks & Rubiales, 2002). Pre-formed barriers also include chemical compounds present on the leaf surface, for example, antimicrobial enzymes and secondary metabolites, which might inhibit spore germination (Heath, 1997a; Heath, 1997b; Morrisey & Osbourn, 1999;...
Figure 1: Proposed stages of nonhost resistance against *Alternaria brassicae*. This schematic representation depicts the possible events during the interaction of *Alternaria brassicae* with the host (A) and non-host plants (B), which is divided into four different phases. The NHR towards *A. brassicae* may involve the pre-formed and inducible defense barrier at different phases. I. The first phase involves the deposition and germination of *A. brassicae* spores on the leaf surface. Nonhost plants have a low frequency of spore deposition on the leaf surface due to the differences in their canopy structure (continued on next page...).
compared to the host plant. Moreover, spore germination might be inhibited due to the presence of pre-formed antimicrobial compounds on the leaf surface and irregular germination of spores due to the absence of host stimuli. II. The second phase involves germ tube development from the spores and the recognition of the stomata. The difference in the leaf surface topography of the nonhost plant compared to the host plant may not allow the proper development of the germ tube into the appressorium. At the next level, even if the appressorium is formed, either stomata are closed by plant or the pathogen may fail to recognize the stomata due to the absence of specific cues, generally derived from the distinct composition and amount of cuticular waxes in the nonhost leaf, which might be entirely different in the case of the host leaf. III. The third phase involves fungal hyphal formation and its development into mycelium. The nonhost plant may inhibit hyphal formation and its development into mycelium by secreting inducible chemicals by pre-formed antimicrobial compounds or by limiting nutrient availability in the intercellular spaces. IV. The fourth phase involves the penetration of the fungal pathogen into the plant cell wall and its subsequent entry inside the plant cell. The nonhost plant may inhibit the penetration of the fungal hyphae into the plant cell wall by inducing the formation of papillae. The defense responses are induced by nonhost plants after recognizing PAMPs or elicitor molecules. This may lead to the production of enzymes and a large amount of phytoalexins by the nonhost plants, which detoxify the toxins produced by the fungal pathogen and restrict fungal growth and multiplication. Note: The text on the right in the figure indicates the NHR mechanisms, and the highlighted blue text indicates the most exploitable NHR mechanisms to develop durable disease-resistant Brassica crops.

Heath, 2000). The ability of A. brassicae spores to germinate on nonhost tomato and wheat plants was tested by McRoberts & Lennard (1996). They found that the spores were able to germinate with the same efficiency on the nonhost and host plants. These spores may have overcome the pre-formed chemical barriers, thereby being able to germinate on the leaf surface. Furthermore, the entry of germinated A. brassicae spores inside the plants is achieved either by targeting the stomata, which is a more predominant mode for entry, or directly through the cuticle (Changsri & Weber, 1960; Tsuneda & Skoropad, 1978). Stomatal recognition by a pathogen requires specific plant topographical cues, which allow germ tube development and eventually facilitate the penetration inside the plant (Hoch et al., 1987; Heath, 1997a; Staples & Hoch, 1997; Niks & Rubiales, 2002). A study on tomato and wheat plants showed that germ tube development of A. brassicae was not affected, but the pathogens failed to enter the stomata of the nonhost plants, unlike in the host plants (McRoberts & Lennard, 1996). It is possible that in the nonhost plants, stomata are not accurately recognized by the pathogen because the surface topography may significantly differ from that of the host leaf. The lack of stomatal recognition by the pathogen may lead to the development of the germ tube, albeit away from the location of the stomatal opening. This sort of disruption in the orientation of the germ tube may not allow the pathogen to enter inside the leaf tissue. Another pre-formed structural barrier includes cuticular waxes, which retard pathogen entry into the leaf tissues (Tewari & Skoropad, 1976; Skoropad & Tewari, 1977; Conn, 1986; Serrano et al., 2015). For example, barley has excessive cuticular wax, consisting of approximately 54% long-chain β-diketones, 20% hydroxylated β-diketones, and other hydrocarbons (Wettstein-Knowles & Netting, 1976; Simpson & Von Wettstein-Knowles, 1980). Together, these form thick cuticular layers that occlude the stomatal features involved in germ tube penetration inside the leaf tissue (Uppalapati et al., 2012). Besides, Alternaria, being a necrotrophic pathogen, sometimes directly enters the host plant through the cuticle by enzymatically digesting cutin waxes by
means of cutinolytic enzymes (Tewari, 1986; Conn & Tewari, 1989). For a few tolerant host species, a high content of cuticular waxes acts as a physical barrier that provides resistance to *A. brassicicola* (Tewari & Skoropad, 1976; Skoropad & Tewari, 1977; Conn & Tewari, 1989). We propound that nonhost plants may have appreciably high amounts of cuticular waxes, which may restrict pathogens from penetrating inside the plant. The penetration attempts by the pathogen on the nonhost plant may trigger inducible defense responses in the plant. The nonhost plant can induce stomatal closure, prevent the entry of *A. brassicicola*, and mount an inducible chemical barrier involving the rapid formation of phytoalexins—antimicrobial compounds inhibiting hyphal development and differentiation (Grayer & Harborne, 1994; Mert-Turk, 2002; Iriti & Faoro, 2007). The differentiation of hyphae is important for penetration through intercellular spaces, and certain nutrients are essential for hyphal development and differentiation (Giri et al., 2013). Thus, in a nonhost plant, the lack of nutrients and presence of antimicrobial compounds in the apoplast may inhibit the development of hyphae into mycelium. Moreover, the defense responses also involve the formation of a structural barrier to arrest fungal growth and penetration in the cell wall (Underwood, 2007; Voigt, 2015). For example, *A. brassicicola* fails to penetrate the cell wall in tomato and wheat plants, and callose-containing papilla formation is observed in the cell wall regions where the pathogen attempts to penetrate such plants (McRoberts & Lennard, 1996). The pathogen also produces nonhost-specific or general toxins that can damage the plant cells, which eventually leads to necrosis (Buchwaldt & Green, 1992; Bains, Tewari & Ayer, 1993; Parada et al., 2007). To avoid this, a nonhost plant may recognize these toxins and employ defense mechanisms to detoxify these toxins (Pedras et al., 2001). Some NHR mechanisms operate at the post-invasion level, which might restrict the pathogen from invading the epidermal cells in the nonhost plant. These defense responses mainly involve ROS accumulation and defense-induced cell death, which do not allow the mycelium to ramify through and between the cells. Eventually, fungal proliferation might be restricted by rapid HR (McRoberts & Lennard, 1996; Gilchrist, 1998).

**Understanding the molecular events in NHR to other *Alternaria* species**

Defense responses are evoked in both host and nonhost infections; however, the NHR involves not only earlier induction but also a robust defense response compared to the host plant. Both *A. brassicicola* and *A. brassicicola* affect almost the same cruciferous crops and generally exhibit similar symptoms (Saharan, Mehta & Meena, 2016; Meena et al., 2016). However, the literature indicates that *Arabidopsis* and *S. alba*, which belong to the same family (Brassicaceae) show NHR to *A. brassicicola* (Thomma et al., 1999; Pedras et al., 2001) but act as hosts for *A. brassicicola* (Hansen & Earle, 1997; Sharma et al., 2002). A recent study showed *Arabidopsis* accession Gre-0 to be highly susceptible to *A. brassicicola* (Mandal, Rajarammohan & Kaur, 2018). Therefore, understanding the NHR mechanisms in these plants against *A. brassicicola* will indirectly illustrate the resistance mechanisms operating during the interaction of *A. brassicicola* with nonhost plants or absent during the interaction with these susceptible host plants. Hence, this section covers the molecular mechanisms of NHR to *A. brassicicola*. 
In A. thaliana and S. alba, early and high-level induction of defense-related genes, namely, pathogenesis-related-1 (PR1), β-1,3 glucanase (PR2), and chitinase (PR3), occurs compared to B. juncea after infection with A. brassicicola (Narusaka et al., 2003; Van Wees et al., 2003; Ghose et al., 2008; Nayanakantha et al., 2016; Mandal, Rajarammohan & Kaur, 2018). Moreover, the expression of PR3, encoding chitinase, remained undetected in B. juncea after pathogen infection, but its expression was highly induced in S. alba (Nayanakantha et al., 2016). However, both A. thaliana and S. alba actively secrete chitinase enzymes, which hydrolyze the fungal cell wall and release chitin fragments (Narusaka et al., 2003; Chatterjee, Mazumder & Basu, 2013). These chitin fragments are recognized by some receptors that activate effective defense responses against the pathogens (Wan et al., 2008). This suggests that the defense-related genes are activated earlier in nonhost plants than in host plants as a part of the NHR strategy, and the chitinase capable of degrading fungal cell walls plays an important role in restricting pathogen growth at early stages of infection. Furthermore, the receptors involved in chitin recognition in Arabidopsis, e.g., CERK 1 or LysM receptor-like kinase (LysM-RLK) and phytosulfokine receptor kinase (PSK), were found to be induced after A. brassicicola infection (Wan et al., 2008; Loivamäki et al., 2010). Wan et al. (2008) also showed that a mutation in the LysM-RLK gene inhibits the induction of all chitin-responsive defense genes and compromises the NHR to A. brassicicola. Similarly, Arabidopsis homologs for RLK receptors were identified in S. alba, and their expression was highly induced after infection with A. brassicicola; the genes were downregulated in B. juncea (Ghose et al., 2008). This indicates that chitin digestion from the fungal cell wall and its perception by the plant may play an essential role during defense signaling in nonhost plants. The NHR response involves the stimulation of a signal transduction cascade after the perception of a pathogen by the plant cell, which initiates the activation of protein kinases and members of MAP kinases and subsequently leads to defense gene activation in nonhost plants (Lawrence et al., 2008). MAPK6 expression was found to be highly induced for a longer duration in S. alba plants infected with A. brassicicola, but it was downregulated in B. juncea (Taj et al., 2010). These findings lead us to speculate that MAPK6 might be involved in imparting NHR and targeting many downstream components that are involved in effective defense response.

Salicylic acid (SA), jasmonic acid (JA), and abscisic acid (ABA) are the key modulators involved in phytohormone signaling during plant defense. JA-mediated signaling is predominantly involved in general defense against necrotrophic pathogens (Glazebrook, 2005). In Arabidopsis, JA-mediated activation of the defense response occurs against A. brassicicola (Thomma et al., 1998; Van Wees et al., 2003), while S. alba challenged with the same pathogen induces ABA-mediated and JA-mediated defense responses (Mazumder et al., 2013). In contrast, susceptible B. juncea plants induce SA-mediated defense signaling pathways (Thomma et al., 1998; Van Wees et al., 2003; Mazumder et al., 2013). It is likely that there is a significant amount of crosstalk between phytohormones and a convergence of two or more signaling pathways, which play a role in deciding whether disease progression will occur or the defense pathways will overcome the pathogen. Considering these studies.
together, we surmise that early signaling events in nonhost plants has a major role in imparting NHR to *A. brassicicola*.

**Possible strategies for developing durable Alternaria blight-resistant *Brassica* crops by using genomic tools**

Based on a literature survey on intercropping systems, we have shortlisted the nonhost plants for *A. brassicae* (Table 1) that can be chosen for dissecting the NHR mechanisms. As discussed earlier, the NHR to *A. brassicae* may involve four different phases based on fungal invasion and penetration events (Fig. 1). Prior to any specific study, it is important to classify the nonhost plants based on these four phases by using pathological assays. This will provide a clear idea about whether the NHR operates at the pre-invasion or post-invasion level. Now, we outline the five major strategies for working towards the ultimate goal of developing durable Alternaria blight-resistant *Brassica* crops.

**Inter or intra-specific introgression of candidate genes from nonhost plants to *Brassica* crops**

The first strategy involves the identification of genes conferring NHR by utilizing conventional or molecular breeding approaches, followed by the introgression of candidate genes from nonhost plants to *Brassica* crops. This can be achieved in two ways. First, where possible, interspecific crosses of the host with a nonhost may be performed to transfer resistance traits from the nonhost plant to a host plant. The unwanted background in the progeny can be removed by crossing it with wild-type host plants and selecting for the targeted traits only. Then, the progeny can be screened for response to *A. brassicae*. Plants showing resistance to the pathogen can be subsequently subjected to QTL mapping to identify the genes involved in NHR. Once the genes are identified, they can be successfully introgressed into the desired varieties to provide resistance to *A. brassicae*. The major limitation with this approach, however, is that the progeny generally suffers from sterility and abnormal segregation, which can hinder the identification of genes involved in resistance. To our knowledge, the only available example of an interspecific cross made between a host and nonhost plant is that of *Lactuca* and *Bremia* (*Jeuken et al.*, 2008).

The second approach is to cross two different varieties of nonhost plants (intraspecific cross: completely resistant genotype × moderately resistant genotype) that differ in the degree of resistance towards *A. brassicae*. This will allow the inheritance of NHR-specific QTL of the resistant donor into the moderately susceptible genotypes of the same species of the nonhost plant (*Jafary et al.*, 2008; *Niks & Marcel*, 2009). The progeny obtained can be similarly studied as described above for identifying the genes involved in NHR and transferring them into target crops. However, if the nonhost plant does not have significant variation in the degree of resistance towards the pathogen, the inheritance of NHR-specific genes might be difficult (Fig. 2).

**Microarray-based gene identification and transfer**

The second strategy is to perform microarray analysis after infecting the nonhost plant (e.g., chickpea or barley) with *A. brassicae* and then identify the candidate genes from the list of upregulated or downregulated genes. If a homologous gene is identified, mutant or
Figure 2 Schematic representation of the possible strategies for exploiting the NHR mechanisms to develop durable blight-resistant *Brassica* crops. The selection of a suitable nonhost plant is essential for exploiting NHR mechanisms. NHR may operate at two different layers, i.e., pre-invasive or post-invasive layers. Once the exploitable NHR layer is identified in the selected nonhost plant, several strategies can be suitably utilized for identifying the genes and pathways involved in the NHR mechanisms. The first approach involves global transcriptome profiling specific to *A. brassicaceae*-induced responses from the nonhost plant to identify genes playing a role in two different layers of NHR. The data can be enriched by using proper controls, e.g., non-adapted pathogens, and the genes exclusively involved in the NHR can be selected. The second approach involves the use of functional genomic tools, e.g., VIGS, TIGS, and T-DNA/transposon insertional mutagenesis, for identifying the genes.
overexpression lines can be generated for the identified genes in the susceptible model plant. Alternatively, the candidate genes can be cloned from the nonhost plant and expressed in the susceptible model plant. These transgenic plants can then be tested for their response to *A. brassicae* infection. Finally, the genes imparting NHR to *A. brassicae* can be successfully transferred to *Brassica* crops.

**Gene silencing**

The third strategy to identify genes underlying NHR involves gene silencing in the nonhost plant. Virus-induced gene silencing (VIGS)-based forward genetic screening can be performed in nonhost plants that have established VIGS protocols and cDNA libraries to identify the genes involved in NHR to *A. brassicae*. For example, for wheat and barley, the *Barley stripe mosaic virus* (BSMV)-based vector and a well-developed VIGS protocol for gene silencing are available (*Ma et al., 2012*). Such a strategy will be useful in identifying important genes involved in NHR by screening a VIGS population for its response to *A. brassicae*. Furthermore, high-throughput transient expression and gene silencing methods, e.g., transient-induced gene silencing (TIGS), have been used in a nonhost plant such as barley. This technique is based on the transformation of leaf epidermal cells by the biolistic method (*Douchkov et al., 2014a; Douchkov et al., 2014b*). TIGS constructs may be made for the genes that are upregulated in the nonhost plant after *A. brassicae* infection. These TIGS constructs can be bombarded into leaf epidermal cells of the nonhost plant to induce gene silencing. After bombardment, leaf discs can be infected with *A. brassicae* and screened for the response. TIGS will allow the identification of potentially important candidate genes involved in NHR to *A. brassicae*. The genes identified using any of these functional genomic methods can be successfully transferred to *Brassica* crops to ensure resistance to *A. brassicae*.

**Identification and use of model plants as genetic resources**

The fourth strategy is to work with model plants that have well-developed molecular and genetic resources and a large population of mutants as nonhosts for *A. brassicae*. For example, *Brachypodium distachyon* can be directly utilized for identifying genes and dissecting the pathways involved in NHR (*An et al., 2016*). If the mutants are not readily available, then with the help of well-developed genetic tools, transposon or T-DNA insertional mutagenesis can be attempted in the model nonhost plant. The mutant population can be screened for the response to *A. brassicae* to identify the candidate genes involved in NHR. These candidate genes can then be successfully transferred to *Brassica* crops for durable resistance to *A. brassicae*. *Arabidopsis* is a well-established, easy-to-manipulate model plant with ample genetic resources. Another approach can be screening the response of all available *Arabidopsis* accessions to *A. brassicae*. This may help identify *Arabidopsis* accessions imparting resistance to *A. brassicae* (*Rajarammohan et al., 2017; Mandal, Rajarammohan & Kaur, 2018*).

**Literature mining and targeted functional genomics**

The fifth strategy is solely based on literature mining and targeted functional genomics to dissect the different layers of NHR involved in pre- and post-invasive defense responses.
The available literature provides hints about possible candidates specifically involved in pre- and post-invasive defenses. This will provide the opportunity to directly start studying these candidates for their role in NHR. Another possibility is that the post-invasive defenses may play a more important role in restricting the fungal pathogen rather than pre-invasive defenses against *A. brassicae*, which can be determined by performing pathological assays. In such cases, we can use mutants with mutations in the genes important for pre-invasion defense to identify the genes involved in post-invasion defenses. For example, the mutant of the barley gene *Ror1*, a homolog of the *Arabidopsis* gene *Pen1*, is important for pre-invasive defense against many nonhost pathogens, including rust pathogens (*Collins et al., 2003; Trujillo et al., 2004*). This mutant line can be used for identifying genes involved in the post-invasion defense response. The response of a *ror1* mutant line should be tested towards *A. brassicae* to ascertain whether this gene is involved in pre-invasive defense against this pathogen. Subsequently, the NHR-specific candidate genes can be identified by transcriptomic analysis in wild-type barley after *A. brassicae* infection. Gene silencing approaches can be applied to silence the candidate genes in the *ror1* mutant line for identifying their role in post-invasion immunity by screening them against *A. brassicae*. Furthermore, validation experiments can be carried out by constructing double mutants of candidate genes and *Ror1* in barley. The double mutants compromising NHR and showing a higher degree of susceptibility to *A. brassicae* compared to the single mutant *ror1* can be likely candidates that contribute to NHR at the post-invasion level. The important genes identified can be further transferred, or their homologs can be overexpressed in the susceptible *Brassica* crops to test their resistance to *A. brassicae*.

**Conclusion and future perspectives**

Selection from within the existing germplasm and the utilization of these sources to incorporate resistance in cultivated varieties through conventional breeding techniques have embodied the basic approaches for generating disease-resistant varieties. However, the exploitation of resistance mechanisms from host germplasm sources has shown limited success in the development of disease-resistant crop varieties. Meanwhile, when genes are introduced from related genera or families, the durability of resistance might be questionable, particularly when a single gene regulating one pathway for defense is employed. Therefore, it is essential to conduct parallel research based on the exploitation of NHR sources or genes against the pathogen over the course of developing varieties resistant to Alternaria blight. Compared to other forms of resistance, NHR governs defense responses to a broad range of pathogens. It is a stable, durable, and robust form of resistance that relies on multiple defense components. Sustainable and broad-spectrum resistance under field conditions makes NHR a promising resource for crop improvement. We speculate that the strategies proposed for exploiting NHR by applying the latest molecular biology techniques involving transcriptomic approaches and genomic tools such as VIGS, TIGS, and mutagenesis will be useful in identifying novel sources of NHR and developing *Brassica* crops with durable disease resistance. The utilization of model plants can be pivotal in identifying the genes and deciphering the pathways operational during NHR. A model plant can serve as a promising tool to enable plant breeders to transfer
the knowledge obtained from it into crop plants for developing disease-resistant crops. However, the utilization of NHR sources for effective crop improvement is beset by several limitations, the major one being that one sole NHR mechanism cannot be exploited to combat all the pathogens of a crop. NHR is a more complex form of defense due to the involvement of multiple pathways; hence, it is less feasible in crop breeding applications. Furthermore, pathogens having a rapid evolutionary rate in nature tend to overcome the NHR mechanisms operating in the nonhost plants.

**ACKNOWLEDGEMENTS**

Authors thank Dr. Muthamilarasan Mehanathan for critical reading of the manuscript.

**ADDITIONAL INFORMATION AND DECLARATIONS**

**Funding**
Muthappa Senthil-Kumar and Priyadarshini Bhorali labs are funded by Department of Biotechnology-North East Region (DBT-NER) twinning project (BT/PR15998/NER/95/145/2015). Urooj Fatima was funded by a DBT-SRF fellowship (DBT/2013/NIPGR/68). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**Grant Disclosures**
The following grant information was disclosed by the authors:
Department of Biotechnology-North East Region (DBT-NER) twinning project: BT/PR15998/NER/95/145/2015.
DBT-SRF fellowship: DBT/2013/NIPGR/68.

**Competing Interests**
The authors declare there are no competing interests.

**Author Contributions**
- Urooj Fatima analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
- Priyadarshini Bhorali and Sudarshana Borah prepared figures and/or tables, authored and reviewed drafts of the paper, approved the final draft.
- Muthappa Senthil-Kumar conceived and designed the experiments, contributed reagents/materials/analysis tools, authored or reviewed drafts of the paper, approved the final draft.

**Data Availability**
The following information was supplied regarding data availability:
This is a review article. Table S1 provides information on resistant sources. No other raw data are available for the article.
Supplemental Information
Supplemental information for this article can be found online at http://dx.doi.org/10.7717/peerj.7486#supplemental-information.

REFERENCES

Agarwal A, Garg GK, Devi S, Mishra DP, Singh US. 1997. Ultrastructural changes in Brassica leaves caused by Alternaria brassicae and destruxin B. Journal of Plant Biochemistry and Biotechnology 6:25–28 DOI 10.1007/BF03263004.

Agnihotri A, Gupta K, Prem D, Sarkar G, Mehra VS, Zargar SM. 2009. Genetic enhancement in rapeseed- mustard for quality and disease resistance through in vitro techniques. In: Proceedings of 16th Australian research assembly on Brassicas, Ballarat, Australia. 28.

Agnihotri A, Lakshmikumaran MS, Jagannathan V, Shivanna KR. 1991. Wide hybridization for improvement in cultivated Brassicas. Acta Horticulture (ISHS) 289:213–214.

Agrios GN. 2005. Plant pathology. Fifth edition. Burlington: Elsevier Academic Press, 948.

Ahlawat IPS, Sharma RP. 2002. Agronomic terminology. 50. New Delhi: Indian Society of Agronomy, Division of Agronomy, Indian Agriculture Research Institute, 132.

Ahlawat IPS, Gangaiah B, Ompal S. 2005. Production potential of chickpea (Cicer arietinum) based intercropping systems under irrigated conditions. Indian Journal of Agronomy 50:27–30.

An T, Cai Y, Zhao S, Zhou J, Song B, Bux H, Qi X. 2016. Brachypodium distachyon T-DNA insertion lines: a model pathosystem to study nonhost resistance to wheat stripe rust. Scientific Reports 6:25510 DOI 10.1038/srep25510.

Aneja JK, Agnihotri A. 2013. Alternaria blight of oilseed Brassicas: epidemiology and disease control strategies with special reference to use of biotechnological approaches for attaining host resistance. Journal of Oilseed Brassica 4:1–10.

Ao E, Saud KTR. 2016. Production potential of rapeseed (Brassica rapa var. dichotoma)—based intercropping systems under rain fed conditions. Journal of Oilseed Brassica 1:91–97.

Awasthi RP, Kolte SJ. 1994. Epidemiological factors in relation to development and prediction of Alternaria blight of rapeseed and mustard. Indian Phytopathology 47:395–399.

Bains PS, Tewari JP, Ayer WA. 1993. A note on phytotoxicity of homodestruxin B—a compound produced by Alternaria brassicae. Phytoprotection 74(3):157–160.

Buchwaldt L, Green H. 1992. Phytotoxicity of destruxin B and its possible role in the pathogenesis of Alternaria brassicae. Plant Pathology 41:55–63 DOI 10.1111/j.1365-3059.1992.tb02316.x.

Changsri W, Weber GF. 1960. Studies of Alternaria spp. pathogenic on Cruciferae. Phytopathology 50:9.
Chatterjee M, Mazumder M, Basu D. 2013. Functional analysis of the promoter of a glycosyl hydrolase gene induced in resistant *Sinapis alba* by *Alternaria brassicicola*. *Phytopathology* 103:841–850 DOI 10.1094/PHYTO-11-12-0303-R.

Chavan V, Bhargava S, Kamble A. 2013. Temporal modulation of oxidant and antioxidative responses in *Brassica carinata* during β-aminobutyric acid-induced resistance against *Alternaria brassicaceae*. *Physiology and Molecular Plant Pathology* 83:35–39 DOI 10.1016/j.pmpp.2013.03.002.

Chevre AM, Eber F, Margale E, Kerlan MC. 1994. Comparison of somatic and sexual *Brassica napus* *Sinapis alba* hybrids and their progeny by cytogenetic studies and molecular characterization. *Genome* 37:367–371 DOI 10.1139/g94-052.

Cho Y, Davis JW, Kim KH, Wang J, Sun QH, Cramer CB, Lawrence CB. 2006. A high throughput targeted gene disruption method for *Alternaria brassicicola* functional genomics using linear minimal element (LME) constructs. *Molecular Plant-Microbe Interaction* 19:7–15 DOI 10.1094/MPMI-19-0007.

Chupp C, Sherf AF. 1960. *Vegetable diseases and their control*. New York: The Ronald Press Communication, 693.

Collins NC, Thordal-Christensen H, Lipka V, Bau S, Kombrink E, Qiu JL. 2003. SNARE-protein-mediated disease resistance at the plant cell wall. *Nature* 425:973–977 DOI 10.1038/nature02076.

Conn KL. 1986. Leaf epicuticular wax of canola: ultrastructure, chemistry and interaction with *Alternaria brassicaceae*. M. S.C. thesis, University of Alberta, Edmonton, 159.

Conn KL, Jejelowa OA, Tewari JP, Bains PS. 1991. *Camelina sativa* phytoalexins (Calmexin and methoxy camalexin) provide resistance against *Alternaria brassicaceae* [Abstract 274]. *Canadian Journal of Plant Pathology* 13.

Conn KL, Tewari JP. 1989. Interactions of *Alternaria brassicaceae* conidia with leaf epicuticular wax of canola. *Mycology Research* 93:240–242 DOI 10.1016/S0953-7562(89)80126-1.

Conn KL, Tewari JP, Awasthi RP. 1990. A disease assessment key for *Alternaria blackspot* in rapeseed and mustard. *Canadian Plant Disease Survey* 70:19–22.

Conn KL, Tewari JP, Dahiya JS. 1988. Resistance to *Alternaria brassicaceae* and phytoalexin-elicitation in rapeseed and other crucifers. *Plant Science* 55:21–25 DOI 10.1016/0168-9452(88)90037-4.

Conn KL, Tewari JP, Hadziyev D. 1984. The role of epicuticular wax in canola in resistance to *Alternaria brassicaceae* [Abstract 851]. *Phytopathology* 74.

Douchkov D, Baum T, Ihlow A, Schweizer P, Seiffert U. 2014b. Microphenomics for interactions of barley with fungal pathogens. In: Tuberosa R, Graner A, Frison E, eds. *Genomics of plant genetic resources*. 2nd edition. Heidelberg: Springer, 123–148.

Douchkov D, Luck S, Johrde A, Nowara D, Himmelbach A, Rajaraman J, Schweizer P. 2014a. Discovery of genes affecting resistance of barley to adapted and non-adapted powdery mildew fungi. *Genome Biology* 15(12):Article 518 DOI 10.1186/gb-2014-15-1-r1.
Dzurilla M, Kutschy P, Tewari JP, Ruzinsky M, Senvicky S, Kovacik V. 1998. Synthesis and antifungal activity of new indolypthiazinone derivatives. *Collection Czechoslovakia Chemical Communication* 63:94–102 DOI 10.1135/ccce19980094.

Ebrahimi E, Kaul HP, Neugschwandtner RW, Dabbagh Mohammadi Nassab A. 2016. Productivity of wheat (*Triticum aestivum* L.) intercropped with rapeseed (*Brassica napus* L). *Canadian Journal of Plant Science* 97:557–568.

Ellis J. 2006. Insights into nonhost disease resistance: can they assist disease control in agriculture? *The Plant Cell* 18:523–528 DOI 10.1105/tpc.105.040584.

Fonseca JP, Mysore KS. 2018. Genes involved in nonhost disease resistance as a key to engineer durable resistance in crops. *Plant Science* 279:108–116.

Gangasaran G, Giri G. 1985. Intercropping of mustard with chickpea, lentil and barley in drylands. *Indian Journal of Agronomy* 30:244–250.

Ghose K, Dey S, Barton H, Loake GJ, Basu D. 2008. Differential profiling of selected defence-related genes induced on challenge with *Alternaria brassicicola* in resistant white mustard and their comparative expression pattern in susceptible India mustard. *Molecular Plant Pathology* 9:763–775 DOI 10.1111/j.1364-3703.2008.00497.x.

Gilchrist DG. 1998. Programmed cell death in plant disease: the purpose and promise of cellular suicide. *Annual Review of Phytopathology* 36:393–414 DOI 10.1146/annurev.phyto.36.1.393.

Giri P, Taj G, Meena PD, Kumar A. 2013. Microscopic study of *Alternaria brassicae* infection processes in *Brassica juncea* cultivars by drop plus agarose method. *African Journal of Microbiology Research* 7:4284–4290.

Glazebrook J. 2005. Contrasting mechanisms of defense against biotrophic and necrotrophic pathogens. *Annual Review of Phytopathology* 43:205–227 DOI 10.1146/annurev.phyto.43.040204.135923.

Grayer RJ, Harborne JJ. 1994. A survey of antifungal compounds from higher plants. *Phytochemistry* 37:19–42 DOI 10.1016/0031-9422(94)85005-4.

Gupta SK, Gupta PP, Kaushik CD. 1995. Changes in leaf peroxidase, polyphenols oxidase, catalase and total phenols due to *Alternaria* leaf blight in *Brassica* species. *Indian Journal of Mycology and Plant Pathology* 25:175–180.

Gupta SK, Gupta PP, Yadav TP, Kaushik CD. 1990. Metabolic changes in mustard due to *Alternaria* leaf blight. *Indian Phytopathology* 43:64–69.

Gupta SK, Kaushik CD. 2002. Metabolic changes in mustard leaf and silique wall due to the infection of *Alternaria* blight (*Alternaria brassicae*). *Cruciferae Newsletter* 24:85–86.

Hansen LN, Earle ED. 1997. Somatic hybrids between *Brassica oleracea* and *Sinapis alba* L. with resistance to *Alternaria brassicae* (Berk.) Sacc. *Theory of Applied Genetics* 94:1078–1085 DOI 10.1007/s001220050518.

Heath MC. 1997a. Evolution of plant resistance and susceptibility to fungal parasites. In: Carroll GC, Tudzynski P, eds. *The Mycota V, part B, plant relationships*. Berlin: Springer, 257–276.

Heath MC. 1997b. Signalling between pathogenic rust fungi and resistant or susceptible host plants. *Annual of Botany* 80:713–720 DOI 10.1006/anbo.1997.0507.
Heath MC. 2000. Nonhost resistance and nonspecific plant defenses. Current Opinion of Plant Biology 3:315–319 DOI 10.1016/S1369-5266(00)00087-X.

Hoch HC, Staples RC, Whitehead B, Comeau J, Wolf ED. 1987. Signaling for growth orientation and cell differentiation by surface topography in uromyces. Science 235:1659–1662 DOI 10.1126/science.235.4796.1659.

Humpherson-Jones FM, Maude RB. 1982. Studies on the epidemiology of Alternaria brassicicola in Brassica oleracea seed production crops. Annals of Applied Biology 100(1):61–71.

Humpherson-Jones FM, Phelps K. 1989. Climatic factors influencing spore production in Alternaria brassicae and A. brassicicola. Annual of Applied Biology 114:449–458 DOI 10.1111/j.1744-7348.1989.tb03360.x.

Iriti M, Faoro F. 2007. Review on innate and specific immunity in plants and animals. Mycopathology 164:57–64 DOI 10.1007/s11046-007-9026-7.

Ishiga Y, Uppalapati SR, Gill US, Huhman D, Tang Y, Mysore KS. 2015. Transcriptomic and metabolomic analyses identify a role for chlorophyll catabolism and phytoalexin during Medicago nonhost resistance against Asian soybean rust. Scientific Reports 5:13061 DOI 10.1038/srep13061.

Jafary H, Albertazzi G, Marcel TC, Niks RE. 2008. High diversity of genes for non-host resistance of barley to heterologous rust fungi. Genetics 178:2327–2339 DOI 10.1534/genetics.107.077552.

Jejelowo OA, Conn KL, Tewari JP. 1991. Relationship between conidial concentration, germling growth, and phytoalexin production by Camelina sativa leaves inoculated with Alternaria brassicae. Mycology Research 95:928–934 DOI 10.1016/S0953-7562(09)80089-0.

Jeromela AM, Mikić AM, Vujeć S, Ćupina B, Krstić D, Dimitrijević A, Vasiljević S, Mihailović V, Cvejić S, Miladinović D. 2017. Potential of legume–Brassica intercrops for forage production and green Manure: encouragements from a temperate southeast european environment. Frontiers in Plant Science 8:312.

Jeukjen MJW, Pelgrom K, Stam P, Lindhout P. 2008. Efficient QTL detection for nonhost resistance in wild lettuce: backcross inbred lines versus F2 population. Theoretical and Applied Genetics 116:845–857 DOI 10.1007/s00122-008-0718-2.

Kadian AK, Saharan GS. 1983. Symptomatology, host range and assessment of yield losses due to Alternaria brassicae infection in rapeseed and mustard. Indian Journal of Mycology and Plant Pathology 13:319–323.

Kanrar S, Venkateswari JC, Kirt PB, Chopra VL. 2002. Transgenic expression of hevein, the rubber tree lectin, in Indian mustard confers protection against Alternaria brassicae. Plant Science 162:441–448 DOI 10.1016/S0168-9452(01)00588-X.

Kohmoto K, Otani H, Tsuge T. 1995. Alternaria alternata pathogens. Vol. 2 Eukaryotes, Turkey: Oxford Pergamon, 51–63.

Kolte SJ. 1985. Diseases of annual edible oilseed crops. Vol. II, rapeseed-mustard and sesame diseases. Boca Raton: CRC Press Inc, 135.
Kolte SJ. 1996. Diseases. In: Chopra VL, Prakash S, eds. Oilseeds and vegetable Brassicas: An Indian perspective. 294. New Delhi: Oxford and IBH Publishing Co. Pvt. Ltd, 184–207.

Kolte SJ. 2002. In: Rai M, Singh H, Hegde DM, eds. Diseases and their management in oilseed crops, new paradigm in oilseeds and oils: research and development needs. Hyderabad: Indian Society of Oilseeds Research, 244–252.

Krishnia SK, Saharan GS, Singh D. 2000. Genetics of alternaria blight resistance in inter and intraspecific crosses of Brassica juncea and B. carinata. Annual of Biology 16:212–216.

Kumar D, Maurya N, Bharati YK, Kumar A, Kumar K, Srivastava K, Chand G, Singh SK, Mishra RK, Kumar A. 2014. Alternaria blight of oilseed Brassicas: a comprehensive review. African Journal of Microbiology Research 8:2816–2829 DOI 10.5897/AJMR2013.6434.

Lawrence CB, Mitchell TK, Craven KD, Cho YR, Cramer RA, Kim KH. 2008. At death’s door: Alternaria pathogenicity mechanisms. Plant Pathology Journal 24:101–111 DOI 10.5423/PPJ.2008.24.2.101.

Lee HA, Lee HY, Seo E, Lee J, Kim SB, Oh S, Choi E, Choi E, Lee SE, Choi D. 2017. Current understandings of plant nonhost resistance. Molecular Plant-Microbe Interactions 30:5–15 DOI 10.1094/MPMI-10-16-0213-CR.

Lipka V, Dittgen J, Bednarek P, Bhat R, Wiermer M, Stein M, Landtaq J, Brandt W, Rosahl S, Scheel D, Llorente F, Molina A, Parker J, Somerville S, Schulze-Lefert P. 2005. Pre-and postinvasion defenses both contribute to nonhost resistance in Arabidopsis. Science 310:180–1183.

Loivamaki M, Stuhrwohldt N, Deeken R, Steffens B, Hirnrich R, Sauter M. 2010. A role for PSK signaling in wounding and microbial interactions in Arabidopsis. Physiologia Plantarum 139(4):348–357 DOI 10.1111/j.1399-3054.2010.01371.x.

Ma M, Yan Y, Huang L, Chen M, Zhao H. 2012. Virus-induced gene-silencing in wheat spikes and grains and its application in functional analysis of HMW-GS-encoding genes. BMC Plant Biology 12:1 DOI 10.1186/1471-2229-12-1.

Mandal S, Rajarammohan S, Kaur J. 2018. Alternaria brassicaceae interactions with the model Brassicaceae member Arabidopsis thaliana closely resembles those with Mustard (Brassica juncea). Physiology and Molecular Biology of Plants 24:51–59 DOI 10.1007/s12298-017-0486-z.

Mathur B, Chand L. 1991. Effect of Alternaria brassicaceae toxin on photosynthesis and photorespiration in Brassica cultivars. Crop Research 4:146–151.

Maude RB, Humpherson-Jones FM. 1980. Studies on the seed-borne phases of dark leaf spot (Alternaria brassicicola) and grey leaf spot (Alternaria brassicae) of brassicas. Annual of Biology 95:331–319.

Mazumder M, Das S, Saha U, Chatterjee M, Bannerjee K, Basu D. 2013. Salicylic acid-mediated establishment of the compatibility between Alternaria brassicicola and Brassica juncea is mitigated by abscisic acid in Sinapis alba. Plant Physiology and Biochemistry 70:43–51 DOI 10.1016/j.plaphy.2013.04.025.
McRoberts N, Lennard JH. 1996. Pathogen behaviour and plant cell reactions in interactions between Alternaria species and leaves of host and nonhost plants. *Plant Pathology* 45(4):742–752.

Medhi BN. 1985. Intermating and mutagenesis for release of genetic variability in Indian mustard (*Brassica juncea* L Czern and Coss). PhD thesis, Punjab Agriculture University, Ludhiana.

Meena PD, Awasthi RP, Chattopadhyay C, Kolte SJ, Kumar A. 2010. Alternaria blight: a chronic disease in rapeseed-mustard. *Journal of Oilseed Brassica* 1:1–11.

Meena PD, Awasthi RP, Chattopadhyay C, Kolte SJ, Kumar A. 2016. Alternaria blight: a chronic disease in rapeseed-mustard. *Journal of Oilseed Brassica* 1:1–11.

Meena PD, Rani A, Meena R, Sharma P, Gupta R, Chowdappa P. 2012. Aggressiveness, diversity and distribution of *Alternaria brassicicola* isolates infecting oilseed *Brassica* in India. *African Journal of Microbiology Research* 6:5249–5258.

Mert-Turk F. 2002. Phytoalexins: defence or just a response to stress? *Journal of Cell and Molecular Biology* 1:1–6.

Mohanty A, Chhrungu B, Verma N, Shivanna KR. 2009. Broadening the genetic base of crop *Brassicas* by production of new intergeneric hybrid. *Czech Journal of Genetics Plant Breeding* 45:117–122 DOI 10.17221/35/2009-CJGPB.

Mondal KK, Bhattacharya RC, Koundal KR, Chatterjee SC. 2007. Transgenic Indian mustard (*Brassica juncea*) expressing tomato glucanase leads to arrested growth of *Alternaria brassicicola*. *Plant Cell Reports* 26:247–252 DOI 10.1007/s00299-006-0241-3.

Mondal KK, Chatterjee SC, Viswakarma N, Bhattacharya RC, Grover A. 2003. Chitinase mediated inhibitory activity of *Brassicas* transgenic on growth of *Alternaria brassicicola*. *Current Microbiology* 47:171–173 DOI 10.1007/s00284-002-3980-6.

Morrisey JP, Osbourn AE. 1999. Fungal resistance to plant antibiotics as a mechanisms of pathogenesis. *Microbiology and Molecular Biology Reviews* 63:708–724.

Mysore KS, Ryu CM. 2004. Nonhost resistance; how much do we know? *Trends in Plant Science* 9:97–104.

Narusaka Y, Narusaka M, Seki M, Ishida J, Nakashima M, Kamiya A, Enju A, Sakurai T, Satoh M, Kobayashi M. 2003. The cDNA microarray analysis using an *Arabidopsis pad3* mutant reveals the expression profiles and classification of genes induced by *Alternaria brassicicola* attack. *Plant Cell and Physiology* 44:377–387 DOI 10.1093/pcp/pcg050.

Nayanakantha NMC, Rawat S, Ali S, Grover A. 2016. Differential expression of defense-related genes in *Sinapis alba* and *Brassica juncea* upon the infection of *Alternaria brassicicola*. *Tropical and Agricultural Research* 27(2):123–136.

Niks RE, Marcel TC. 2009. Nonhost and basal resistance: how to explain specificity? *New Phytologist* 182:817–828 DOI 10.1111/j.1469-8137.2009.02849.x.

Niks RE, Rubiales D. 2002. Potentially durable resistance mechanisms in plants to specialized fungal pathogens. *Euphytica* 124:201–216 DOI 10.1023/A:1015634617334.

Nishimura S, Kohmoto K. 1983. Host-specific toxins and chemical structures from *Alternaria* species. *Annual Review of Phytopathology* 21:87–116 DOI 10.1146/annurev.py.21.090183.000511.
Nurnberger T, Lipka V. 2005. Non-host resistance in plants: new insights in an old phenomenon. *Molecular Plant Pathology* 6:335–345 DOI 10.1111/j.1364-3703.2005.00279.x.

Parada RY, Oka K, Yamagishi D, Kodama M, Otani H. 2007. Destruxin B produced by *Alternaria brassicae* does not induce accessibility of host plants to fungal invasion. *Physiology and Molecular Plant Pathology* 71:48–54 DOI 10.1016/j.pmpp.2007.10.003.

Pedras MSC, Smith KC. 1997. Sinalexin, a phytoalexin from white mustard elicited by destruxin B and *Alternaria brassicae*. *Phytochemistry* 46:833–837 DOI 10.1016/S0031-9422(97)00362-2.

Pedras MSC, Zaharia IL, Gai Y, Zhou Y, Ward DE. 2001. In planta sequential hydroxylation and glycosylation of a fungal phytotoxin: avoiding cell death and overcoming the fungal invader. *Proceedings of the National Academy of Sciences of the United States of America* 98:747–752 DOI 10.1073/pnas.98.2.747.

Prasad R, Saxena D, Chandra S. 2003. Yield losses by *Alternaria* blight in promising genotypes of Indian mustard. *Indian Phytopathology* 56:205–206.

Primard C, Vedel F, Mathieu C, Pelletier G, Chevre AM. 1988. Interspecific somatic hybridization between *Brassica napus* and *Brassica hirta* (*Sinapis alba* L.). *Theory of Applied Genetics* 75:546–552 DOI 10.1007/BF00289119.

Rajarammohan S, Kumar A, Gupta V, Pental D, Pradhan AK, Kaur J. 2017. Genetic architecture of resistance to *Alternaria brassicae* in *Arabidopsis thaliana*: QTL mapping reveals two major resistance-conferring loci. *Frontiers in Plant Science* 8:260.

Saharan GS. 1992. *Management of rapeseed and mustard diseases*. vol. 1. India: Advances of Oilseed Research Science Publication, 152–533.

Saharan GS, Mehta N, Meena PD. 2016. The disease. In: *Alternaria diseases of crucifers: biology, ecology and disease management*. Singapore: Springer, 17–51.

Senthil-Kumar M, Mysore KS. 2013. Nonhost resistance against bacterial pathogens: retrospectives and prospects. *Annual Review of Phytopathology* 51:407–427 DOI 10.1146/annurev-phyto-082712-102319.

Serrano M, Coluccia F, MartínaTorres FLH, Métraux JP. 2015. The cuticle and plant defense to pathogens. *Frontiers in Plant Science* 5:274.

Sharma G, Kumar DV, Haque A, Bhat SR, Shyam P, Chopra VL. 2002. *Brassica* coenospecies: a rich reservoir for genetic resistance to leaf spot caused by *Alternaria brassicae*. *Euphytica* 125:411–419 DOI 10.1023/A:1016050631673.

Sharma TR, Singh BM. 1992. Transfer of resistance to *Alternaria brassicae* in *Brassica juncea* through interspecific hybridization among brassicas. *Journal of Genetics and Breeding* 46:373–378.

Shekhawat K, Rathore SS, Premi OP, Kandpal BK, Chauhan JS. 2012. Advances in agronomic management of Indian mustard (*Brassica juncea* (L.) Czernj. Cosson): an overview. *International Journal of Agronomy* 2012:1–14.

Shyam SY, David L, Philip C. 2007. *Lentil an ancient crop for modern times*. vol. 472. Dordrecht: Springer, 259.
Simpson D, Von Wettstein-Knowles P. 1980. Structure of epicuticular waxes on spikes and leaf sheaths of barley as revealed by a direct platinum replica technique. *Carlsberg Research Communications* 45:465–481 DOI 10.1007/BF02932920.

Singh DN, Singh NK, Srivastava S. 1999. Biochemical and morphological characters in relation to Alternaria blight resistance in rape-seed mustard. *Annual Agriculture Research* 20:472–477.

Singh RK, Kumar H, Singh AK. 2010. *Brassica* based intercropping systems—a review. *Agriculture Review* 3:253–266.

Singh VS, Kothari K, Tripathi HN. 1986. Studies on intercropping in sugarcane in central Uttar Pardesh. *Indian Sugarcane Journal* 35:559–562.

Singh V, Kumar K, Bhajan R, Singh PK, Singh RB. 2009. Identification of resistance sources against Alternaria blight and white rust in Indian mustard. *Journal of Oilseeds Research* 26:435–436.

Skoropad WP, Tewari JP. 1977. Field evaluation of the epicuticular wax in rapeseed and mustard in resistance to *Alternaria brassicae*. *Canadian Journal of Plant Science* 57:1001–1003 DOI 10.4141/cjps77-146.

Staples RC, Hoch HC. 1997. Physical and chemical cues for spore germination and appressorium formation by fungal pathogens. In: *Plant relationships*. Berlin Heidelberg: Springer, 27–40.

Subudhi PK, Raut RN. 1994. White rust resistance and its association with parental species type and leaf waxiness in *B. juncea* L. Czern and Coss. X *B. napus* L. Crosses under the action of EDTA and Gamma ray. *Euphytica* 74:1–7 DOI 10.1007/BF00033760.

Suman A, Lal M, Singh AK, Gaur A. 2006. Microbial biomass turnover in Indian subtropical soils under different sugarcane intercropping systems. *Agronomy Journal* 98:698–704 DOI 10.2134/agronj2005.0173.

Taj G, Agarwal P, Grant M, Kumar A. 2010. MAPK machinery in plants: recognition and response to different stresses through multiple signal transduction pathways. *Plant Signaling and Behavior* 5:1370–1378 DOI 10.4161/psb.5.11.13020.

Tewari JP. 1983. Cellular alterations in the black spot of rapeseed caused by *Alternaria brassicae*. *Phytopathology* 73:831–831.

Tewari JP. 1986. Subcuticular growth of *Alternaria brassicae* in rapeseed. *Canadian Journal of Botany* 64:1227–1231 DOI 10.1139/b86-168.

Tewari JP. 1991. Structural and biochemical bases of black spot disease of crucifers. *Advances in Structural Biology* 1:325–349.

Tewari JP, Conn KL, Dahiya JS. 1987. Resistance to *Alternaria brassicae* in crucifer. In: *Proceedings of 7th International Rapeseed Congress, Poznan*, vol. 48. 1085–1089.

Tewari JP, Conn KL, Dahiya JS. 1988. Dynamic response of rapeseed and other crucifers to *Alternaria brassicae* and *Rhizoctonia solani*. In: *Proceedings of 5th international congress of plant pathology, Kyoto*. 242.

Tewari JP, Skoropad WP. 1976. Relationship between epicuticular wax and blackspot caused by *Alternaria brassicae* in three lines of rapeseed. *Canadian Journal of Plant Science* 56:781–785 DOI 10.4141/cjps76-127.

Fatima et al. (2019), *PeerJ*, DOI 10.7717/peerj.7486
Thomma BPHJ. 2003. *Alternaria* spp. from general saprophyte to specific parasite. *Molecular Plant Pathology* 4:225–236 DOI 10.1046/j.1364-3703.2003.00173.x.

Thomma BPHJ, Eggermont K, Penninckx IAMA, Mauch-Mani B, Vogelsang R, Cammue BPA, Broekaert WF. 1998. Separate jasmonate-dependent and salicylate-dependent defense-response pathways in *Arabidopsis* are essential for resistance to distinct microbial pathogens. *Proceedings of the National Academy of Sciences of the United States of America* 95:15107–15111 DOI 10.1073/pnas.95.25.15107.

Thomma BP, Eggermont K, Tierens KF, Broekaert WF. 1999. Requirement of functional ethylene-insensitive 2 gene for efficient resistance of *Arabidopsis* to infection by *Botrytis cinerea*. *Plant Physiology* 121:1093–1101 DOI 10.1104/pp.121.4.1093.

Tewari KP, Tomar RKS, Mishra GL, Raghu JS. 1992. Intercropping of mustard with gram and lentil. *Journal of Oilseeds Research* 9:248–252.

Toor SS, Singh S, Garch AIS, Kapur ML, Singh A. 2000. Gobhi sarson as an intercrop in Autumn Sugarcane for higher returns. *Intensive Agriculture* 38:29–30.

Tripathi NN, Kaushik CD, Yadav TP, Yadav AK. 1978. Studies on the inheritance of Alternaria blight resistance in raya. *Indian Phytopathology* 31:127.

Tripathi NN, Kaushik CD, Yadav TP, Yadav AK. 1980. Alternaria leaf spot resistance in raya. *Haryana Agriculture University Journal of Research* 10:166–168.

Trujillo M, Troeger M, Niks RE, Kogel KH, Hu ckelhoven R. 2004. Mechanistic and genetic overlap of barley host and non-host resistance to *Blumeria graminis*. *Molecular Plant Pathology* 5:389–96 DOI 10.1111/j.1364-3703.2004.00238.x.

Tsuneda A, Skoropad WP. 1977. Formation of microsclerotia and chlamydospores from conidia of *Alternaria brassicae*. *Canadian Journal of Botany* 55:1276–1281 DOI 10.1139/b77-148.

Tsuneda A, Skoropad WP. 1978. Phylloplane fungal flora of rapeseed. *Transactions of the British Mycological Society* 70:329–333 DOI 10.1016/S0007-1536(78)80130-2.

Underwood W. 2007. The plant cell wall: a dynamic barrier against pathogen invasion. *Frontiers in Plant Science* 3:67.

Uppalapati SR, Ishiga Y, Doraisswamy V, Bedair M, Mittal S, Chen JH, Nakashima J, Tang YH, Tadege M, Ratet P, Chen RJ, Schultheiss H, Mysore KS. 2012. Loss of abaxial leaf epicuticular wax in *Medicago truncatula* irg1/palm1 mutants results in reduced spore differentiation of anthracnose and nonhost rust pathogens. *The Plant Cell* 24:353–370 DOI 10.1105/tpc.111.093104.

Van Wees SC, Chang HS, Zhu T, Glazebrook J. 2003. Characterization of the early response of *Arabidopsis* to *Alternaria brassicicola* infection using expression profiling. *Plant Physiology* 132:606–617 DOI 10.1104/pp.103.022186.

Verma PR, Saharan GS. 1994. Monograph on Alternaria diseases of crucifers. In: *saskatoon research centre technical bulletin*. 6th edition. Saskatoon: Agriculture and Agri-Food Canada.

Verma VD, Rai B. 1980. Note on induced mutagenesis for spotting out the sources of resistance to Alternaria leaf spot in Indian mustard. *Indian Journal of Agriculture Science* 50:278–280.
Vishwanath, Kolte SJ, Singh MP, Awasthi RP. 1999. Induction of resistance in mustard (*Brassica juncea*) against Alternaria black spot with an avirulent *Alternaria brassicace* isolate-D. *European Journal of Plant Pathology* **105**:217–220. DOI 10.1023/A:1008717323002.

Voigt CA. 2015. Callose-mediated resistance to pathogenic intruders in plant defense-related papillae. *Frontier in Plant Science* **5**:168.

Wan J, Zhang XC, Neece D, Ramonell KM, Clough S, Kim SY, Stacey G. 2008. A LysM receptor-like kinase plays a critical role in chitin signaling and fungal resistance in *Arabidopsis*. *The Plant Cell* **20**:471–481. DOI 10.1105/tpc.107.056754.

Wettstein-Knowles P, Netting AG. 1976. Composition of epicuticular waxes on barley spikes. *Carlsberg Research Communication* **41**:225–235. DOI 10.1007/BF02906259.

Yang Z, Rogers LM, Song Y, Guo W, Kolattukudy PE. 2005. Homoserine and asparagine are host signals that trigger in plant expression of a pathogenesis gene in *Nectria haematococca*. *Proceedings of the National Academy of Sciences of the United States of America* **102**:4197–4202. DOI 10.1073/pnas.0500312102.

Zhang FL, Xu JB, Takahata Y. 1996. Inheritance of resistance to black leaf spot (*Alternaria brassicae*) in Chinese cabbage. *Cruciferae Newsletter* **18**:134–135.

Zhang F, Xu J, Yan H, Li M. 1997. Inheritance of resistance to black leaf spot in Chinese cabbage. *Acta Agriculturae Boreali-Sinica* **12**:115–119.