The potential of plant proteins as antifungal agents for agricultural applications

Tiffany Chiu\textsuperscript{a}, Theo Poucet\textsuperscript{b}, Yanran Li\textsuperscript{b,*}

\textsuperscript{a} Graduate Program in Genetics, Genomics, And Bioinformatics, 1140 Bachelor Hall, University of California Riverside, California, 92521, USA
\textsuperscript{b} Department of Chemical and Environmental Engineering, University of California, Riverside, CA, 92521, USA

\textbf{A B S T R A C T}

Fungal pathogens induce a variety of diseases in both plants and post-harvest food crops, resulting in significant crop losses for the agricultural industry. Although the usage of chemical-based fungicides is the common way to control these diseases, they damage the environment, have the potential to harm human and animal life, and may lead to resistant fungal strains. Accordingly, there is an urgent need for diverse and effective agricultural fungicides that are environmentally- and eco-friendly. Plants have evolved various mechanisms in their innate immune system to defend against fungal pathogens, including soluble proteins secreted from plants with antifungal activities. These proteins can inhibit fungal growth and infection through a variety of mechanisms while exhibiting diverse functionality in addition to antifungal activity. In this mini review, we summarize and discuss the potential of using plant antifungal proteins for future agricultural applications from the perspective of bioengineering and biotechnology.

1. Introduction

The management of plant diseases is one of the top priorities of the agricultural industry due to the major economic and biosecurity threats that result from plant pathogens [1]. Among the pathogen cortege that crops are afflicted by, fungal infections pose one of the largest risks to food production [2]. Devastating crop failures due to these pathogens, such as the historical and infamous Irish Potato Famine [3] and contemporary issues of rice blast and wheat rust threaten food security and result in major economic losses [4]. The development of fungicides has undoubtedly eased the burden of diminished food security through a reduction in crop failures by successfully controlling fungal diseases. Chemical fungicides, made from either organic or inorganic chemicals, remain as the primary treatment towards most fungal pathogens [5]. However, chemical fungicides have long been documented for their adverse effects on both the environment and animal health [6], and harvested crops must meet strict criteria to ensure chemical residues are found at safe levels for consumptions [7]. Though conventional fungicides have made positive strides in food security and agricultural disease control, the risks they carry need to be addressed and alternative methods of fungal control should be considered.

One common alternative to conventional fungicides is the usage of genetically modified (GM) crops. Transgenic technology has led to the development of crops with desirable traits, such as improved flavor [8], increased yield [9], and superior disease resistance [10] compared to non-modified crops. Notably, the use of transgenic crops permits for a significant reduction in the quantity of phytosanitary product applied to the field [11]. However, the public is often apprehensive about GMO safety and has difficulty accepting genetically modified crops [12]. For example, some consumers believe that GM crops carry more risks than benefits and are willing to pay a premium for foods labeled as non-GMO [13]. Likewise, since 2001, the EU has placed a de facto moratorium on approvals of GMOs [14]. Another major concern includes the potential that transgenic crops could damage the ecosystem in unpredictable ways. GMOs can invade ecosystems due to an increase in stress tolerance, causing wild plants to become weeds through horizontal gene transfer [15], or produce toxic substances to pests that may affect nontarget organisms [16]. Recently, increases in pest resistance towards GM crops have also posed problems to the durability of current transgenic crops [17].

Thus, it is necessary to seek alternative antifungal agent candidates that can be applied exogenously as conventional fungicides. These alternative candidates should be environmentally friendly and potentially have fewer negative health impacts on animals than conventional fungicides if applied exogenously. Plants have evolved diverse mechanisms to defend against fungal infections, as summarized in Fig. 1, with one important route utilizing the secretion of proteins to delay fungal infection or inhibit fungal growth. These plant antifungal proteins are...
promising candidates since they are biodegradable, generally nontoxic to humans and antagonistic microorganisms, and most importantly, have evolved for millions of years to combat phytopathogenic fungi with a narrow target range [18]. In this mini review, we summarize and discuss plant defensive proteins that are promising candidates for the development of future antifungal agents for agricultural applications (as summarized in Table 1).

2. Pathogenesis related proteins

Pathogenesis related (PR) proteins are a group of low molecular weight plant proteins involved in mitigating both biotic and abiotic stresses [19], and are often involved in triggering systemic acquired resistance in plants [20]. There are 16 main groups of PRs (PR-1 to PR-16), with each group classified based on different molecular and physiological properties. These proteins are often pathogen specific and involved in the transcriptional activation of plant defenses [21]. Here, we will focus on some of the most promising candidates for the development of antifungal agents for agricultural applications: PR-3, PR-5, PR-6, and PR-12.

2.1. Chitinases (PR-3)

One of the best known and most studied plant antifungal proteins is chitinase, which belongs to PR-3 [22]. Chitinases are strongly induced when the host plants are under attack from pathogens and function as defense molecules against fungal infection [23]. These proteins inhibit fungal growth by lysing hyphal tips in fungi and break down chitin into defense molecules against fungal infection [23]. These proteins inhibit fungal growth by lysing hyphal tips in fungi and break down chitin into defense molecules against fungal infection [23]. These proteins inhibit fungal growth by lysing hyphal tips in fungi and break down chitin into defense molecules against fungal infection [23]. These proteins inhibit fungal growth by lysing hyphal tips in fungi and break down chitin into defense molecules against fungal infection [23]. These proteins inhibit fungal growth by lysing hyphal tips in fungi and break down chitin into defense molecules against fungal infection [23]. These proteins inhibit fungal growth by lysing hyphal tips in fungi and break down chitin into defense molecules against fungal infection [23]. These proteins inhibit fungal growth by lysing hyphal tips in fungi and break down chitin into defense molecules against fungal infection [23]. These proteins inhibit fungal growth by lysing hyphal tips in fungi and break down chitin into defense molecules against fungal infection [23]. These proteins inhibit fungal growth by lysing hyphal tips in fungi and break down chitin into defense molecules against fungal infection [23]. These proteins inhibit fungal growth by lysing hyphal tips in fungi and break down chitin into defense molecules against fungal infection [23]. These proteins inhibit fungal growth by lysing hyphal tips in fungi and break down chitin into defense molecules against fungal infection [23].

Fig. 1. Mode of actions of secreted plant antifungal proteins with potential agricultural applications. 1) Secreted antifungal proteins reduce fungal hyphae growth by compromising the fungal cell wall and membrane integrity, leading to potential cytoplasmic leakages [165]. 2) Antifungal protein activity generates residues considered as microbe-associated molecular pattern molecules that can be recognized by plant receptors to stimulate plant immune response [166]. 3) Plant antifungal proteins, upon interacting with the target, directly stimulate plant immune response [167]. 4) Plant secreted proteins protect antifungal proteins from cleavage by fungal protease [168]. 5) Inhibition of fungal protease by plant secreted inhibitors [169]. 6) Inhibition of fungal cell wall hydrolase by plant secreted inhibitors [170]. 7) Spore degradation or reduction of germination rate by secreted plant antifungal proteins [171]. 8) Small secreted peptides enhance the efficacy of plant defense [172].
Table 1
Summary of antifungal proteins of potential to be developed into alternative fungi control agents for agricultural applications.

| Protein Class              | Protein                          | Exogenous Application Inhibition | Antifungal Mechanism                                                                 | Ref           |
|----------------------------|----------------------------------|----------------------------------|--------------------------------------------------------------------------------------|---------------|
| Pathogenesis Related Proteins | Chitinase                        | Alternaria solani, Aspergillus niger, Botrytis cinerea, Collectotrichum falacatum, Fusarium sp, Pestalotia theae, Rhizoctonia solani, Sphaeroteca humuli, Trichoderma sp | Degradation of chitin via hydrolysis of the N-acetylglucosamine polymer, Lysing of fungal hyphal tips | [25,27,36,43,154–158] |
|                            |                                  |                                  | • Unknown                                                                            | [50]          |
|                            |                                  |                                  | • Ion channel inhibition                                                             | [54]          |
|                            |                                  |                                  | • Interacts with fungal cytoplasmic agents                                           | [56,57]       |
|                            |                                  |                                  | • Disrupts plasmic membrane integrity                                               |               |
|                            |                                  |                                  | Disrupts plasmic membrane integrity                                                 | [58]          |
|                            |                                  |                                  | Inhibition of cell wall barriers via signal transduction pathway                     | [62,69,71,159] |
|                            |                                  |                                  | Reduction of pathogen toxicity towards host                                          |               |
|                            |                                  |                                  | Disruption of fungal cell walls, fungal hyphael, and spore germination              |               |
|                            |                                  |                                  | Hydrolyse β-1,3-glucans                                                             |               |
|                            |                                  |                                  | Fungal membrane permealization                                                      |               |
| Protease Inhibitors        | Potide-G                          | Candida albicans, Rhizoctonia solani, Botrytis cinerea, Fusarium oxysporum, Fusarium solani | Competitive and noncompetitive inhibition of serine, aspartic, and cysteine proteases | [76]          |
|                            | Potato Protease Inhibitor 1 and II| Botrytis cinerea, Fusarium oxysporum, Fusarium solani | Chymotrypsin and serine protease inhibition                                             | [75,78,160]   |
|                            | Bowman-Birk Protease Inhibitor    | Fusarium graminearum, Fusarium graminearum, Fusarium moniliforme, Fusarium oxysporum, Fusarium roseum, Kabatiella seae, Phytophthora infestans, Phytophthora parasitica, Sclerotinia sclerotiorum, Stemcapella maydis, Trichoderma longibrachiatum, Verticillium dahliae | Pro tease serine protease inhibitor, Noncompetitive inhibition of trypsin and chymotrypsin | [92,93,161,162] |
| Antimicrobial Peptides     | Prosystemin                       | Botrytis cinerea | Induces protease inhibitors                                                          | [104,105]     |
|                            | StSN1                             | Botrytis cinerea, Fusarium sp. | Amplifies defense signaling process for wounded plants, Mechanism unknown            | [111,163]     |
|                            | Puroindoline A and B              | Alternaria brassicola, Ascochyta pisi, Fusarium culmorum, Fusarium graminearum, Magnaporthe grisea, Rhizoctonia solani, Verticillium dahliae | Induces membrane instability                                                          | [116,164]     |
| DUP26-Containing Proteins  | Ginkobilobin2                      | Candida albicans, Fusarium oxysporum, Trichoderma reesei | Binds sugar motifs on hyphal surface                                                  | [122,123]     |
|                            | AFP1/ AFP2                        | Ustilago maydis, None tested      | Binds to sugar motifs on hyphal surface                                               | [124]         |
|                            | VdCRRR                            | Bipolaris sorokiniana, Rhizoctonia cerealis | Stabilize/protect chitinases from fungal proteases                                    | [125]         |
|                            | TaCRRR                            | Bipolaris sorokiniana, Rhizoctonia cerealis | Inhibited mycelia growth, activation of pathogenesis-related genes                   | [126]         |
| Leucine Rich Repeat Protein| PvPGP2                            | Aspergillus niger, Botrytis cinerea | Competitive and noncompetitive inhibition                                             | [144]         |
| S Albumin and 2S Albumin Orthologs | 2S Albumin                        | Aspergillus flavus, Aspergillus fumigatus, Candida albicans, Fusarium oxysporum, Phanerochaete chrysosporium, Trichoderma harisianum | Mechanism unknown                                                                   | [146,150–152] |
both plant extraction and heterologous expression from microbial factors, and numerous studies documenting its antifungal efficacy, chitinase is a promising candidate to develop as an antifungal agent. However, compared to the number of available chitinase studies that use transgenic plants, the exogenous application of chitinases on infected plants remain to be more thoroughly investigated.

### 2.2. Defensins (PR-12)

Defensins belong to group PR-12 [45], and exhibit broad-spectrum activities against different biotic agents including pathogenic fungi [46]. They are named due to the structural and functional similarities to insect and mammalian defensins [47]. Plant defensins are constitutively expressed in the extracellular space of most vegetative and reproductive plant tissues [48] and can be specifically induced under pathogen stress condition [47,49]. Typically, defensins are small soluble cationic proteins, 45–54 amino acid residues in size, exhibiting eight conserved cysteine residues (C1 to C8) with a conserved spacing pattern, and the tertiary structure is supported by at least four disulfide bonds [46]. Defensins remain stable both under extreme temperatures (as high as 90 °C) and very acidic conditions (pH as low as 1) [45]. For instance, NRBAP, a defensin-like protein purified from Phaseolus vulgaris beans, retained its antifungal activity against Mucor racemius up to 100 °C, and in the pH range of 1–13 [50].

Defensins can interact with a significant diversity of biological targets (e.g., proteases [51], protein synthetic machinery [52], α-amylases in insects [53], and ion channels in fungi [54]). One common mechanism that defensins often adopt to inhibit fungal growth is through the disruption of cell plasma membranes. Plant defensins are usually positively charged proteins and interact with anionic moieties in the membrane, such as glycoproteins, sialoproteins, or phosphoproteins [49]. The defensins cover the target membranes until it reaches a concentration threshold, and then disrupts the membrane integrity by affecting the bilayer curvature [55]. One study provided evidence that NaD1 from Nicotiana alata, which displays antifungal activity against several agronomically important filamentous fungi [56], was able to bind to phospholipids phosphatidic acid [57]. To estimate the effect of the total net charge of defensins on the antifungal activity, a mutagenesis analysis was performed on Rs-AFP2 from radish, and the interaction between the defensins and membrane lipids was improved when the net charge of the protein increased [58]. Plant defensin antifungal activity may not be restricted to targeting the membrane of pathogenic fungi. Indeed, the exogenous application of NaD1 is also associated with the entrance of the protein into fungal intracellular space, resulting in granulation of the cytoplasm and cell death [56]. This suggests that plant defensins could also interact with fungal intracellular targets and possibly with DNA, as already demonstrated by ostrich β-defensins, where E. coli growth was inhibited in assays due to interactions between peptides and cytoplasmic targets that curbed DNA, RNA, and protein synthesis [59]. The diversity of antifungal mechanisms and effectiveness of defensins against a wide range of pathogens implies the potential of this protein family as a promising resource for fighting plant pathogens.

### 2.3. Thaumatin-like proteins (PR-5)

Thaumatin-like proteins (TLP) belong to PR-5 family [60]. TLPs are named so due to their structural similarity to thaumatin, a sweet-tasting, non-toxic protein that was first discovered from the fruit of the tropical plant Thaumatococcus danielli [61]. TLPs exhibit a broad range of biological activities, including antifungal activity. Different TLPs inhibit fungal growth through different mechanisms, including but not limited to disrupting fungal membrane [62], inhibiting fungal enzymes such as xylanase [63], inducing apoptosis by binding to specific fungal membrane receptors [64], and hydrolyzing β-1,3-glucans [65]. Osmotin and osmotin-like proteins are among the most studied TLPs of antifungal activity [66]. Osmotin and orthologs have been shown to exhibit broad-spectrum antifungal inhibitory effects [67]. Overexpression of osmotin in transgenic plants delayed disease symptoms from fungal pathogens [68]. Osmotin isolated from tobacco cell suspensions can inhibit the hyphal growth of numerous pathogenic fungi in vitro, including species from Bipolaris, Collectorichum, Fusarium, Kabatiella, Phytophthora, and Trichoderma [69]; and another osmotin-like protein from Solanum nigrum and overexpressed in E. coli can inhibit the growth of phytopathogenic Fusarium solani f. sp. glycines, Macrophomina phaseolina, and Collectorichum glaesporioides, and Collectorichum gossypii var. cephalosporioides at the concentration between 0.1 μg/μL to 0.3 μg/μL [70]. Additionally, an osmotin-like protein from Solanum nigrum L. var indica was shown to inhibit fungal spore germination and permeabilize fungal hyphae in vitro. This protein is also stable and retains its antifungal activity at temperatures as high as 75 °C for 30 min and pH 3–8 [71]. Further functional exploration of TLPs under various stress conditions for in planta assays will be necessary before its development into a reliable antifungal tool [72].

### 2.4. Protease inhibitors (PR-6)

Plant protease inhibitors (PIs), also called PR-6, are important proteins involved in many plant biological processes, including seed germination, protease-related house-keeping functions, and defense against biotic and abiotic stresses [73]. PIs are normally found in amply quantities in seeds and tubers, and plants in the Solanaceae family generally have exceptionally high levels of PIs [74], including some that can be promising candidates of antifungal agents. For instance, potatoes encode several PIs ranging from 4.1 to 39 kDa that exhibit broad-spectrum antifungal activities [75]. Potide-G, a Kunitz-type PI isolated from potato tubers of size 5.5 kDa, inhibits pathogenic fungi Candida albicans and Rhizoctonia solani in vitro even when heated to 70 °C for 20 min, and also exhibits antiviral and antibacterial activities [76]. Similarly, the potato protease inhibitors I and II (PPI-I and PPI-II) can inhibit the growth of various fungi, including B. cinerea [77], Fusarium solani, and Fusarium oxysporum [75]. Both PPI-I and II are heat stable, which can maintain their ability to inhibit F. solani and F. oxysporum growth in vitro at high temperatures as high as 100 °C [78]. PPI-I and II are also nontoxic, as they have been previously been utilized in human clinical trials for appetite control [79]. The extraction of bioactive PPIs from potatoes is laborious and of low yields [80]. They have also been heterologously expressed in Saccharomyces cerevisiae, yet the antifungal activity of the purified protein was not examined [81]. A more economic production method is needed to enable the development of PPIs as antifungal agents for agriculture applications.

Another PII of interest is the Bowman-Birk protease inhibitor (BBI), which is typically under 20 kDa [82,83], contains seven conserved disulfide bonds, and inhibits trypsin and chymotrypsin, which are common enzymes pathogenic fungi utilize when infecting plants [84]. The BBI gene is induced during plant immune responses and overexpression of this gene in plants confers improved disease resistance against both insect and fungal pathogens [85]. BBIs from the legume (Fabaceae) or cereal (Poaceae) family have a double or single inhibitory loop respectively [86], and synthetic peptides that contain only the disulfide-linked, 9-residue long loop have shown to retain their trypsin and chymotrypsin inhibitory activity [87]. This short, truncated form of the protein may be of interest for the development of antifungal agents of smaller molecular weight for easier production and higher stability, compared with larger protein agents. Aside from the small size, BBI is thermostable with the ability to withstand 100 °C for 10 min, tolerates a wide pH range from 1.6 to 8.0, is not allergic, and is approved by the FDA for human consumption [86]. Additionally, unlike some other candidates to be engineered as antifungal agent, BBI has passed phase II human clinical trials and is highly unlikely to be toxic, especially given its prevalence in soy products [89]. BBIs have already been successfully utilized as an exogenously applied antifungal agent in vitro. One study identified that a BBI-type trypsin-chymotrypsin inhibitor purified from broad beans can...
inhibit the growth of *B. cinerea*, *F. oxysporum*, and *M. arachidicola* at a dose as low as 60 μg per plate [90]. Plant BBs have often been isolated from a variety of seeds such as those from *Vigna mungo* [91], *Cajanus cajan* [92], and *Clitoria fragilis* [93] and have been tested for their insecticidal properties. Rice BB has also been expressed in *E. coli* and retained the inhibitory activity. However, the titer is relatively low at 20 mg/L, likely due to the presence of the disulfide bonds that make it prone to forming inclusion bodies [94]. In addition, care should be taken when developing BB as an antifungal agent, as it is a multifunctional PI with a relatively broad activity towards various proteases [95], and may affect beneficial microbiota and fungi in the soil and plants.

3. Antimicrobial proteins

In addition to PRs, antimicrobial peptides (AMPs) are another protein group of interest. AMPs, also known as host defense peptides [96], can be derived from a variety of organisms, including plants, bacteria, and fungi. In plants, AMPs play a role in the plant innate immune system [97]. AMPs that work specifically against fungi are known as antifungal peptides (AFPs) [98], and feature a wide range of functions that are of interest to both pharmaceutical [99] and agricultural industries [100]. Here, we will only discuss AMPs that have shown potential for agricultural applications.

One AFP of interest is tomato systemin, a small peptide of only 18 amino acids long and is involved in inducing the synthesis of PIs in response to plant wounding and damage from herbivores [101]. Research suggests that systemin moves through the plant phloem and helps amplify the signaling process and allows for distal leaves to respond to the wounding [102]. Tomato plants that overexpress pro-systemin, the precursor of systemin, are found to induce high levels of PI proteins even without wounding [103]. Additionally, transgenic plants expressing pro-systemin reduce lesions by at least 50% from *Pythium infestans*, a pathogen that causes late blight [104]. Systemin peptides have been successfully isolated from tomato, sprayed onto grapevine (*Solanum melongena*) and eggplant (*Vitis vinifera*) plants infected with *B. cinerea* [105] at a concentration of 100 μM, and efficiently delayed necrosis of the infected plants.

Snakins are cell wall-associated defensins that are also classified as AMP and believed to play a role in plant growth, signaling, and defense [106]. Snakins isolated from *Solanum tuberosum* (StSN1) are cysteine-rich peptides roughly 6.9 kDa in size [107,108] and the snakins isolated from potato tubers is effective at suppressing both fungal and bacterial growth at concentrations lower than 10 μM [109]. Transgenic potato plants overexpressing the StSN1 gene exhibited reduced symptoms of *R. solani* infections and higher survival rates compared to the wild type plants [110]. Additionally, StSN1 has been shown to be effective in vitro against *B. cinerea* and several *Fusarium* species [111]. However, snakins have been rarely expressed successfully from microbial hosts, often with low yield and insolubility, which hinders in-depth mechanistic characterization of its action towards pathogenic fungi [112].

Another promising group of AFPs are the puroindolines, which are small, amphipathic tryptophan-rich proteins about 13 kDa in size and found only in wheat (*Triticum*) [113]. They are known to inhibit the growth of pathogenic bacteria and fungi with low mammalian toxicity [114], likely through strong binding with microbial membranes and therefore perturbing the membrane integrity [115]. The primary roles of puroindolines include grain hardness and fungal defense. These proteins are believed to protect seeds from fungal attacks during seed development and germination [116]. There are two major puroindolines, Puroindoline A (PINA) and B (PINB) [117]. When the pin genes are overexpressed in transgenic rice, rice displayed significantly enhanced resistance to rice blast caused by *Magnaporthe oryzae* and a reduction in symptoms due to *Rhizoctonia solani* infections [118]. Purified PINA and PINB proteins from wheat were able to inhibit the growth of a variety of pathogenic fungi, including *Alternaria brassicola*, *Ascochyta pisi*, *F. culmorum*, *G. graminearum*, *Magnaporthe grisea*, *R. solani*, and *Verticillium dahlia*. PINA and PINB are stable over a broad range of temperature (70 °C-130 °C) and pH (2.0–12.0) [115,119]. PINs have been heterologously produced in *Pichia pastoris* with a titer up to 14 mg/L taking advantage of puroindoline’s solubility in the detergent Triton X-114 [120]. These various AFPs discussed highlight the potential of using AFPs as antifungal agents for agricultural purposes.

4. DUF26-containing proteins

In the past decade, there has been an increase in interest towards proteins containing domain of unknown function (DUF26) for their capability in fighting plant pathogens and especially fungi [121]. DUF26 is a cysteine rich domain with a conserved C-X8-C-X2-C motif. DUF26-containing proteins are a large, land plant-specific protein family and characteristic of embryophytes [121]. Similarities with fungal lectins suggests DUF26-containing proteins constitute a group of plant carbohydrate-binding proteins able to recognize specific fungal sugar motifs [121].

There are three groups of DUF26-containing proteins: the cysteine-rich receptor-like secreted proteins (CRRSPs), cysteine-rich receptor-like kinase (CRKs) and plasmodesmata-localized proteins (PDLPs). The three DUF26-containing protein groups were all previously associated with antifungal activities. Nevertheless, only CRRSPs remain as good candidates for biotechnological application since CRKs and PDLPs contain transmembrane domains and localize to the membranes. CRRSPs contain a signal peptide followed by one or more DUF26 domains, separated by a variable region [121]. The most well-known CRRSP is Ginkobilobin2 (Gnk2), which was isolated from seeds of *Ginkgo biloba* and able to inhibit the growth of *F. oxysporum*, *T. reesei*, and *C. albicans* [122]. This antifungal activity is likely due to the binding of DUF26 domain with sugar moieties on the fungal cell wall [123]. For instance, Gnk2 interacts specifically with mannann, a yeast cell wall polysaccharide, and mannanse, a building block of mannan, by strictly recognizing the hydroxy group at the C4 position of the monosaccharide. Consistently, two maize CRRSPs (AFP1 and AFP2) have been characterized to interact directly with the hyphal surface of *Ustilago maydis*, and the activity can be rendered by Rsp3, a U. maydis effector covering its surface [124].

In addition to direct binding with fungal cell walls, DUF26-containing proteins from CRRSP family also protect plants using indirect mechanisms. CR1, a secreted apoplastic protein from cotton, and composed of two Cys-rich DUF26 motifs, interacts and protects the antifungal apoplastic chitinase 28 from cleavage by VdSSEP1, a pathogen related protease [125]. Importantly, overexpressing CR1 in heterologous plants such as *Arabidopsis thaliana* and *Nicotiana tabacum* improved plant resistance to *B. cinerea* and *P. parasitica*, respectively. Thus, CR1 could be a good candidate as a co-antifungal agent and simultaneous exogenous application of CR1 and chitinases should be evaluated. Another CRRSP of interest is the currently reported CBM1-interacting protein (OsCBMIP) in rice [126]. Pathogenetic fungi generally use cell wall degrading enzymes (CWDEs) to destruct plant cell walls, and many CWDEs use carbohydrate binding modules (CBMs) to facilitate the access to plant polysaccharides to advance the infection process [127]. OsCBMIP can specifically bind to CBM of several CBM-containing CWDEs including the xylanase MoCel10A of the blast fungus pathogen *Magnaporthe oryzae* and slow down the infection progress. Interestingly, OsCBMIP cannot inhibit the growth of *M. oryzae* and *F. oxysporum* in vitro, and this further indicates that OsCBMIP slows down the infection of pathogenetic fungi through indirect mechanism, here specially, through inhibiting CBM-containing CWDEs [126]. In another study, a transcriptomic analysis of wheat after * Bipolaris sorokiniana* or *Rhizoctonia cerealis* infection reported the induction of a cysteine-rich protein (CBR), TaCRR [128]. When heterologously expressed, this DUF26-containing protein showed a clear antifungal activity. Besides, it was found that silencing TaCRR gene in wheat
significantly decreased the expression of pathogenesis-related genes such as β-1,3-glucanase, defensin or chitinases [128]. Owing to their apoplastic localization and direct or indirect antifungal activities, DUF26-containing proteins from the CRRSP class remain as attractive candidates for the future development of antifungal agents.

5. Other proteins

Polygalacturonase inhibiting proteins (PGIPs) are a family of leucine rich repeat (LRR) proteins found in plant cell walls [129,130] whose primary role is to inhibit polygalacturonases (PGs), enzymes secreted by insects and fungal pathogens that degrade the plant cell walls and leave it vulnerable for infection [131]. Through competitive or noncompetitive inhibition, PGIPs slow the hydrolysis process of PGs [132–135]. Presently, numerous studies show that overexpression of PGIPs in transgenic plants leads to increased fungal resistance. The best-documented PGIP is PGIP2 from Phaseolus vulgaris (PvPGIP2), the common bean [136]. PvPGIP2 has been successfully expressed in transgenic plants, resulting in increased resistance to fungal infections against Alternaria citri, Aspergillus flavus, A. niger, B. cinerea, Claviceps purpurea, and F. graminearum [137–140]. Similarly, expression of PGIP3 from soybeans (Glycine max) in tobacco has been shown to inhibit the growth of pathogenic Sclerotinia sclerotiorum, Fusarium moniliforme, B. aclada, A. niger, Collectothricum atactatum, and F. graminearum [141,142]; and expressing PGIP2 from lime beans (Phaseolus lunatus) in tobacco also delayed growth of Collectothricum lupini, B. cinerea, F. moniliforme, and A. niger [143]. Recently, it is also found that truncated PvPGIP2 with only the optimal docking area retains similar level of inhibitory activities towards PGs from A. niger and B. cinerea to the full-length PvPGIP2 [144]. Yeast strains secreting full-length or truncated PvPGIP2 with the Ots 1 signal peptide were also able to reduce fungal growth and delay sporulation by 1–2 days [144]. Although the function of PGIPs when applied exogenously on plants has not been reported, this group of proteins is still considered as a promising candidate to be developed into an eco-friendly fungal control agent.

Albumins are a major class of water soluble, seed storage proteins that are used as a source of nutrients for plants during germination [145]. Among them, 2S albumins have antifungal capabilities [146], in addition to a variety of activities including anti-cancer, anti-fungal, anti-bacterial, and serine-protease inhibiting properties [147]. These small storage proteins are present in both monocotyledonous and dicotyledonous plant seeds [148] and typically have a disulfide bridge that is thermal-stable at up to 90°C for 50 minutes [149]. For example, pumpkin (Cucurbita sp.) 2S albumin is thermal-stable at up to 90°C, and exhibits inhibitory activity against the fungal pathogen F. oxysporum [147]. Similarly, a crude extract of peanut (Arachis hypogaea) containing 2S albumin was found to inhibit growth of A. flavus [150]; the 2S albumin ortholog from passionfruit (Passifora edulis) could also inhibit the fungal pathogens T. harzianum and F. oxysporum [151], C. musae, and C. lindemuthianum [146]; and the 25S albumin ortholog from Pinnaniju roxburghii (putrin) could inhibit the growth of F. oxysporum, Phanerochaete chrysosporium, C. albicans, Aspergillus fumigatus, and A. flavus. In addition, putrin is stable at up to 50°C and within a pH range from 6 – 8 [152]. On the other hand, 2S albumin from white sesame seeds, oriental mustard, and Brazil nuts can bind to IgG sera, which may trigger an allergic response in humans [145]. Thus, before 2S albumin can be utilized as an exogenously applied antifungal agent, we need to either engineer the protein to eliminate or reduce the allergenicity or modify the application in a manner that avoids either extensive contact or consumption.

6. Conclusion

Chemical-based fungicides are known to be detrimental to the environment and may lead to resistance in pathogenic fungi [153]. Unlike chemical fungicides, the use of exogenously applied natural plant proteins with known antifungal properties can potentially be an eco-friendly and sustainable method for controlling fungal diseases. These natural plant proteins are more socially acceptable, and compared with the production of transgenic plants, are more flexible. Additionally, antifungal plant proteins offer a variety of mechanisms and tools, urgently needed to fight against the rapidly evolving fungal pathogens. As summarized in Table 1, these naturally occurring plant peptides are strong candidates for developing broad-spectrum, fungal-control strategies. One of the biggest hurdles to consider when developing these proteins is lowering the cost of production while enabling mass production. It will be necessary to explore and further optimize microbial factories and protein extraction methods before many of these natural plant proteins can be utilized readily in agricultural industry. Numerous studies showcase the efficacy of these proteins both in vitro and in planta against pathogenic fungi. The potential of using natural plant proteins exogenously to control agricultural fungal diseases remains largely un-tapped and need to be considered when developing future eco- and environmentally-friendly antifungal agents.

Acknowledgments

We thank S. Xu for the valuable feedback in the preparation of the manuscript.

References

[1] Waage JK, Mumford JD. Agricultural biosecurity. Philos. Trans. R. Soc. B Biol. Sci. 2007;363:863–76.
[2] Agrios G. Plant pathology: fifth edition. In: Plant pathology. Fifth; 2004. p. 1–922. 9780808547178.
[3] Rudin R, Donnelly JS. The great Irish potato famine. Can J En Stud 2000. https://doi.org/10.2307/25515536.
[4] Fones HN, et al. Threats to global food security from emerging fungal and oomycete crop pathogens. Nat. Food. 2020;16:1:332–42. 2020.
[5] Morton V, Staub T. A short history of fungicides. APSnet Featur. Artic. 2008. https://doi.org/10.1094/annetfeatur-2008-0308.
[6] Rani L, et al. An extensive review on the consequences of chemical pesticides on human health and environment. J Clean Prod 2021;283:124657.
[7] Lozwickova B, Hrynkov I, Kaczynski P, Jankowska M. Long-Term Investigation and Health Risk Assessment of Multi-class Fungicide Residues in Fruits Estimating acute and chronic exposure of children and adults to chlorpyrifos in fruit and vegetables based on the new, lower toxicology data View project. Artic. Polish J. Environ. Stud. 2016. https://doi.org/10.15254/jjes/611111.
[8] Klee HJ. Improving the flavor of fresh fruits: genomics, biochemistry, and biotechnology. New Phylol 2010. https://doi.org/10.1111/j.1469-8137.2010.03281.x.
[9] Klümper W, Qaim M. A meta-analysis of the impacts of genetically modified crops. PLoS One 2014. https://doi.org/10.1371/journal.pone.0111629.
[10] Borrelli VMG, Brambilla V, Bogowsky P, Marocco A, Lamelbe A. The enhancement of plant disease resistance using crisp/cas9 technology. Front Plant Sci 2018;9:1245.
[11] Dutta SS, et al. Current status and future prospects of research on genetically modified rice: a review. Agric Rev 2016;37.
[12] Hielscher S, Pies I, Valentino V, Chatalova L. Rationalizing the GMO debate: the ordnomic approach to addressing agricultural myths. Int J Environ Res Pub Health. 2016. https://doi.org/10.3390/ijerph130505476.
[13] Zheng Q, Wang H. Do consumers view the genetically modified food labeling systems differently? Contains GMO” Versus “Non-GMO” Labels 2021:376–88. https://doi.org/10.1080/10971475.2021.1890356 54.
[14] Abbas C. EU environmental law: challenges, change and decision-making by maria lee. Mod Law Rev 2006;69.
[15] Warwick SJ, Beckie JJ, Hall LM. Gene flow, invasiveness, and ecological impact of genetically modified crops. Annals of the New York Academy of Sciences; 2009. https://doi.org/10.1111/j.1749-6632.2009.04578.x.
[16] Varrazkas TH, Arvanitoyannis IS, Baltas H. The politics and science behind GMO acceptance. Crit Rev Food Sci Nutr 2007. https://doi.org/10.2307/25515536.
[17] Tabassnik BE, Carriere Y. Surge in insect resistance to transgenic crops and prospects for sustainability. Nat Biotechnol. 2017. https://doi.org/10.1038/nbt.3974.
[18] Zaker M. Natural plant products as eco-friendly fungicides for plant diseases control- A review. Agric For 2016;14:134-41.
[19] Sudisha J, Sharathchandra RG, Amruthesh KN, Kumar A, Shetty HS. Pathogenesis related proteins in plant defense response. Plant Def. Biol. Control. 2012;12: 379–403.
[20] Awasthi, L. P. Applied plant virology : advances, detection, and antiviral strategies.
T. Chiu et al. Synthetic and Systems Biotechnology 7 (2022) 1075–1083

[21] Ali S, et al. Pathogenesis-related proteins and peptides as promising tools for engineering plants with multiple stress tolerance. Microbiol Res. 2018. https://doi.org/10.1016/j.micres.2018.04.008.

[22] Singh A, Isaac Kirubakaran S, Sakhnivet H. Heterologous expression of new antifungal chitinase from wheat. Protein Expr Purif 2007;56:100–9.

[23] Grover A. Plant chitinases: genetic diversity and physiological roles. CRC Crit Rev Plant Sci 2012;31:283–304.

[24] Malik A, Preety. Purification and properties of plant chitinases: a review. J Food Biochem 2019;43:

[25] Baezlen P van, Legendre L, Kan J A L van. Plant defence compounds against botrytis infection. Botytis Biol. Pathol. Control 2007;1:43–61. https://doi.org/10.1007/978-1-4020-2626-3-9.

[26] R D, et al. The Top 10 fungal pathogens in molecular plant pathology. Mol Plant Pathol 2012;14:314–30.

[27] Shrestha CL, Gna, Muthukrishnan S, Mew TW. Chitinase levels in rice cultivars correlate with resistance to the sheath blight pathogen Rhizoctonia solani. Eur J Plant Pathol 2007;120:69–77. 2007 1201.

[28] Molla KA, et al. Understanding sheath blight resistance in rice: the road behind and the road ahead. Biotechnol J 2020;18:895–915.

[29] Hamid R, et al. Chitinases: an update. J Pharm BioAllied Sci 2013;5:21.

[30] Nampoothiri KM, et al. Process optimization for antifungal chitinase production. Process Biochem 2004. https://doi.org/10.1006/sbbi.2003.0282-6.

[31] Brzezinska MS, Jankiewicz U. Production of antifungal chitinase by Aspergillus Niger LOCK 62 and its potential role in the biological control. Curr Microbiol 2004. https://doi.org/10.1007/s00284-012-0208-2.

[32] Muzzarelli RAA, et al. Current views on fungal chitin/chitosan, human chitinases, food preservation, glucans, pectins and inulin: a tribute to Henri Bracconnet, precursor of the carbohydrate polymers science, on the chinin bicentennial. Carbohydr Polym 2012;97:1012–9.

[33] Kasprzewska A. Plant chitinases-regulation and function. Cell Mol Biol Lett 2003;8:

[34] Nagpure A, Choudhary B, Gupta RK. Chitinases: in agriculture and human nutrition. Biotech Lett 2012;34:1065–71.

[35] Hartl L, Zach S, Seid-Selethović B. Chitinases diversity, mechanistic properties and biotechnological potential. Appl Microbiol Biotechnol 2012;93:533–43.

[36] Karamiad T, Sanaka T, Kajihara Y, Yamamoto Y, Koga D. Plant chitinase as a possible biocontrol agent for use instead of chemical fungicides. Biosci Biotechnol Biochem 2003;67.

[37] Bhumian, B. Production and characterization of a thermostable chitinase from a new alkalophilic Bacillus sp. BG-1L.

[38] Landin PGC, et al. Production in Pichia pastoris, antifungal activity and crystal structure of a class I chitinase from copeiva (Vigne ungulifera): insights into sugar binding mode and hydrolytic action. Biochimie 2011;93:89–103.

[39] Dong, X., Zhao, Y., Ran, X., Guo, L. & Zhao, D. G.-O. Overexpression of a new chitinase gene EuCHT2 enhances resistance to Cryptodi cichoracca DC in Tobacco plants. Int. J. Mol. Sci. Artic. doi:10.3390/ijms11122601.

[40] Kurusaki P, Tashiro N, Nishii A. Secretion of chitinase from cultured carrot cells treated with fungal mucoidal w eals. Physiol Mol Plant Pathol 1987;31:211–6.

[41] Hamid R, et al. Chitinases: an update. J Pharm BioAllied Sci 2013;5:521.

[42] Falconer RJ, et al. Thermal stability of thaumatin-like protein, chitinase, and invertebrate isolate of thaumatin-like inhibitors. Insect Sci and Applied ecology. 1994;5:259–66.

[43] Rashel Kahir S, et al. Purification and characterization of a novel chitinase from Trichosanthos dioica seed with antifungal activity. Int J Biol Macromol 2015;74:62–3.

[44] Zhang J, Koppaparar NK, Yan Q, Yang S, Jiang Z. Purification and characterization of a novel chitinase from persimmon (Diospyros kaki) with antifungal activity. Food Chem 2013;138:1225–32.

[45] Edregea A. PATHOGENESIS-RELATED proteins: research progress in the last 15 years. Gen Plant Physiol 2005;51.

[46] Kovalova V, Bakhlejaa I, Kit OY. Plant defensins from a structural perspective, vol. 1:2002. https://doi.org/10.1007/978-1-4020-2626-3-9.

[47] Shrestha CL, Gna, Muthukrishnan S, Mew TW. Chitinase levels in rice cultivars correlate with resistance to the sheath blight pathogen Rhizoctonia solani. Eur J Plant Pathol 2007;120:69–77. 2007 1201.

[48] Molla KA, et al. Understanding sheath blight resistance in rice: the road behind and the road ahead. Biotechnol J 2020;18:895–915.

[49] Hamid R, et al. Chitinases: an update. J Pharm BioAllied Sci 2013;5:21.

[50] Nampoothiri KM, et al. Process optimization for antifungal chitinase production. Process Biochem 2004. https://doi.org/10.1006/sbbi.2003.0282-6.

[51] Kasprzewska A. Plant chitinases-regulation and function. Cell Mol Biol Lett 2003;8:

[52] Kurusaki P, Tashiro N, Nishii A. Secretion of chitinase from cultured carrot cells treated with fungal mucoidal w eals. Physiol Mol Plant Pathol 1987;31:211–6.

[53] Hamid R, et al. Chitinases: an update. J Pharm BioAllied Sci 2013;5:521.

[54] Falconer RJ, et al. Thermal stability of thaumatin-like protein, chitinase, and invertebrate isolate of thaumatin-like inhibitors. Insect Sci and Applied ecology. 1994;5:259–66.

[55] Rashel Kahir S, et al. Purification and characterization of a novel chitinase from Trichosanthos dioica seed with antifungal activity. Int J Biol Macromol 2015;74:62–3.
Giroux, M. J., Sripo, T., Gerhardt, S., & Alfred, RL. Biochemical analysis of the functionality of puroindoline peptides and proteins. J. Biol. Chem. 2017;292:1409-19.

Isaaly, N. et al. Optimization of the wheat puroindoline-a production in Pichia pastoris. Yeast. 2013;29:1409-19.

Vaattovaara, A., et al. Mechanistic insights into the evolution of DUF26-containing proteins in land plants. Commun. Biol. 2019;2:1-18.

Sawano Y, Miyakawa T, Tanokura M, Hatano KJ. Purification, characterization, and molecular gene cloning of an antifungal protein from Ginkgo biloba seeds. Biochem Biophys Res Commun. 2007;388:273-80.

Miyakawa, T. et al. A secreted protein with plant-specific cysteine-rich motif functions as a mannoside-binding lectin that exhibits antifungal activity. Plant Physiol. 2014;166:766-78.

Ma LS, et al. The Ustilago maydis repetitive effector Rpg3 blocks the antifungal activity of mannose-binding proteins. Nat Commun. 2014;5:5192-9.

Han LB, et al. The Cotton Aplastic Protein CR1 stabilizes chitinase 28 to facilitate defense against the fungal pathogen verticillium dahliae. Plant Cell 2019;31:520-36.

Takeda T, et al. Apolipoprotein CMB1 containing proteins bind conserved carbohydrate binding module 1 motifs in fungal hyalodases to counter pathogen invasion. Bioinformatics 2021;37:2474-61. https://doi.org/10.1093/bioinformatics/btaa351.

Takahashi M, et al. Characterization of a celllobiohydrolase (McOsuGA) produced by magnaporthe oryzae. Appl Environ Microbiol. 2010;76:5683-90.

Guo F, Shan Z, Yu J, Xu G, Zhang Z. The cysteine-rich repeat protein tcar1 participates in defense against both rhizoctonia cerealis and bipolaris sorokiniana in wheat. Int J Mol Sci. 2020;21:17.

Di Matteo A, Bonivento D, Tsermoglu D, Federici L, Cervone F. Polygalacturonase-inhibiting protein (PGIP) in plant defence: a structural view. Phytochemistry. 2006. https://doi.org/10.1016/j.phytochem.2005.12.025.

Jones DA, Jones JDG. The role of target proteins and their interactions of fungal polygalacturonases and their plant inhibitors. Phytochemistry. 2006. https://doi.org/10.1016/j.phytochem.2005.12.025.

Jones JDG. Polygalacturonase-inhibiting proteins: in plant innate immunity? Trends Plant Sci. 2006. https://doi.org/10.1016/j.tplants.2005.12.005.

Stover HU, et al. Identification of target amino acids that affect interactions of fungal polygalacturonases and their plant inhibitors. Phytochemistry. 2006. https://doi.org/10.1016/j.phytochem.2005.12.025.

King D, et al. Use of amide exchange mass spectrometry to study conformational changes within the emerging class II homologalacturonanpolygalacturonase-inhibiting protein system. Biochemistry. 2002. https://doi.org/10.1021/bi011291f.

Friediani M, et al. Cytological localization of the PGIP genes in the embryo suspensor cells of Phaseolus vulgaris L. Theor. Appl. Genet. Int. J. Plant Breed. Res. 1993. https://doi.org/10.1007/BF01184925.

Benedetti M, et al. A single amino-acid substitution allows endo-polygalacturonase of Fusarium verticilloides to acquire recognition by PGIP2 from Phaseolus vulgaris. PLoS One 2013. https://doi.org/10.1371/journal. pone.0080610.

Federici L, Di Matteo, A, Fernandez-Rico J, Tsermoglu D, Cervone F. Polygalacturonase-inhibiting proteins in plant immune memory? Trends Plant Sci. 2006. https://doi.org/10.1016/j.tplants.2005.12.005.

Vanaclocha RC, Lima TF, Fernandez-Brum CN, Chalfun-Junior A, Santos JB. Expression and validation of PVP1G genes for resistance to white mold (Sclerotinia sclerotiorum) in common beans (Phaseolus vulgaris L.). Genet Mol Res. 2016. https://doi.org/10.1016/j.gmr.2015.09.042.

Arina F, et al. The bean polygalacturonase-inhibiting protein 2 (PvPGIP2) is highly conserved in common bean (Phaseolus vulgaris L.) germplasm and related species. Theor Appl Genet. 2011;123:1149-59.

Arina F, et al. The bean polygalacturonase-inhibiting protein 2 (PvPGIP2) and PvTAXII-III but not PVPGIP2 and PMEI enhances resistance against Fusarium graminearum, P. varians, and F. avenaceum. World J. Microbiol. Biotechnol. 2019;35:97.

Arina F, Arroyo-Vasquez V, et al. Expression and validation of PVP1G genes for resistance to white mold (Sclerotinia sclerotiorum) in common beans (Phaseolus vulgaris L.). Genet Mol Res. 2016. https://doi.org/10.1016/j.gmr.2015.09.042.

Chiu T, Behari A, Charron JW, Putnam A, Li Y. Exploring the potential of engineered polygalacturonase-inhibiting protein as an ecological, friendly, and nontoxic pest control agent. Biotechnol Bioeng. 2021. https://doi.org/10.1002/biot.2021109.

Moorezij I, Clemeneto A, et al. 25 albumin storage proteins: what makes them food allergens? Open Biochem J. 2008;2:16-28.

Agizzio AP, et al. The antifungal properties of a 25 albumin-homologous protein from passion fruit seeds involve plasma membrane permeabilization and ultrastructural alteration in yeast cells. Plant Sci. 2006;171:515-22.

Tomas PPS, et al. Characterization of antifungal, DNase and antifungal activity of pumpkin 2S albumins. Biochem Biophys Res Commun. 2014;448:349-54.

Souza Candido E, et al. Plant storage proteins with antimicrobial activity: novel insights in plant defense mechanisms. Faseb J 2011;25:3290-305.
Souza PFN. The forgotten 2S albumin proteins: importance, structure, and biotechnological application in agriculture and human health. Int J Biol Macromol 2020;164:4638–49.

Duan XH, Jiang R, Wen YJ, Bin JH. Some 2S albumin from peanut seeds exhibits inhibitory activity against Aspergillus flavus. Plant Physiol Biochem 2013;66:84–90.

Pelegrini PB, et al. An antifungal peptide from passion fruit (Passiflora edulis) seeds with similarities to 2S albumin proteins. Biochim Biophys Acta, Proteins Proteomics 2006;1764:1141–6.

Tomar PFS, et al. Purification, characterization and cloning of a 2S albumin with DNase, RNase and antifungal activities from Putranjiva roxburghii. Appl Biochem Biotechnol 2014;174:471–82.

Wyenandt A, Everts K. Development of a fungicide resistance management guide for vegetable growers in the mid-atlantic states. Mid-atlantic commercial vegetable production recommendations guide view project fungicide resistance management guidelines for vegetable crops grown in the mid-atlantic region. View project 2020. https://doi.org/10.1094/CM-2009-0316-01-MG.

Boller T, Mauch F. Colorimetric assay for chitinase. Methods Enzymol 1988. https://doi.org/10.1016/0076-6879(88)61052-4.

J Jayaraj ZP. Combined expression of chitinase and lipid transfer protein genes in transgenic carrot plants enhances resistance to foliar fungal pathogens. Plant Cell Rep 2007;26:1539–46.

Punja ZK, Zhang YY. Plant chitinases and their roles in resistance to fungal diseases. J Nematol 1993;25(4):526–40.

Singh A, Isaac Kirubakaran S, Sakthivel N. Heterologous expression of new antifungal chitinase from wheat. Protein Expr Purif 2007;56:100–9.

J Z, NK K, Q Y, S Y, Z J. Purification and characterisation of a novel chitinase from persimmon (Diospyros kaki) with antifungal activity. Food Chem 2013;138:1225–32.

Grenier J, Potvin C, Trudel J, Asselin A. Some thaumatin-like proteins hydrolyse polymeric β-1,3-glucans. Plant J 1999;19:473–80.

Hermosa MR, Turra D, Fogliano V, Monte E, Loriito M. Identification and characterization of potato protease inhibitors able to inhibit pathogenicity and growth of Botrytis cinerea. Physiol Mol Plant Pathol 2006;68:138–48.

Chilosi G, et al. Antifungal activity of a Bowman-birk-type trypsin inhibitor from wheat kernel. J Phytopathol 2000;148:477–81.

Katoch R, et al. Cloning, characterization, expression analysis and inhibition studies of a novel gene encoding Bowman-Birk type protease inhibitor from rice bean. Gene 2014;546:342–51.

Segura A, Moreno M, Madueno F, Molina A, García-Olmedo P, Snakin-1, a peptide from potato that is active against plant pathogens. Mol Plant Microbe Interact 1999;12:16–25.

Alfred RL, Palombo EA, Panozzo JF, Bariana H, Bhave M. Stability of puroindoline peptides and effects on wheat rust. World J Microbiol Biotechnol 2013;29:1409–19. 2013 298.

Tanaka S, Kahmann R. Cell wall–associated effectors of plant-colonizing fungi. https://doi.org/10.1080/00275514.2020.1831293 113; 2021. 247-260.

Newman MA, Sundelin T, Nielsen JT, Erbs G. MAMP (microbe-associated molecular pattern) triggered immunity in plants. Front Plant Sci 2013;4:139.

Van J, et al. Plant antifungal proteins and their applications in agriculture. Appl Microbiol Biotechnol 2015;99:4961–81.

Jashni MK, Mehrabi R, Collemare J, Mesarchch, de Wit P.J.GM. The battle in the apoplast: further insights into the roles of proteases and their inhibitors in plant-pathogen interactions. Front Plant Sci 2015;6:584.

Selitrennikoff CP. Antifungal proteins. Appl Environ Microbiol 2001;67:2883–94.

Adams DJ. Fungal cell wall chitinases and glucanases. Microbiology 2004;150:2029–35.

Lanver D, Schweizer G, Tanaka S, Tollot M. Fungal effectors and plant susceptibility. Artic. Annu. Rev. Plant Biol. 2015. https://doi.org/10.1146/annurev-arplant-043014-116623.

Matsubayashi Y. Posttranslationally modified small-peptide signals in plants. 2014. p. 385–413. https://doi.org/10.1146/annurev-arplant-043014-116623.