The vesicular trafficking system component MIN7 is required for minimizing *Fusarium graminearum* infection

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Abstract

Plants have developed intricate defense mechanisms, referred to as innate immunity, to defend themselves against a wide range of pathogens. Plants often respond rapidly to pathogen attack by the synthesis and delivery to the primary infection sites of various antimicrobial compounds, proteins, and small RNA in membrane vesicles. Much of the evidence regarding the importance of vesicular trafficking in plant–pathogen interactions comes from studies involving model plants whereas this process is relatively understudied in crop plants. Here we assessed whether the vesicular trafficking system components previously implicated in immunity in Arabidopsis play a role in the interaction with *Fusarium graminearum*, a fungal pathogen well-known for its ability to cause Fusarium head blight disease in wheat. Among the analysed vesicular trafficking mutants, two independent T-DNA insertion mutants in the *AtMin7* gene displayed a markedly enhanced susceptibility to *F. graminearum*. Earlier studies identified this gene, encoding an ARF-GEF protein, as a target for the HopM1 effector of the bacterial pathogen *Pseudomonas syringae* pv. *tomato*, which destabilizes MIN7 leading to its degradation and weakening host defenses. To test whether this key vesicular trafficking component may also contribute to defense in crop plants, we identified the candidate *TaMin7* genes in wheat and knocked-down their expression through virus-induced gene silencing. Wheat plants in which *TaMin7* genes were silenced displayed significantly more Fusarium head blight disease. This suggests that disruption of MIN7 function in both model and crop plants compromises the trafficking of innate immunity signals or products resulting in hypersusceptibility to various pathogens.

Keywords: Arabidopsis, *AtMin7*, disease resistance, fungal pathogenicity, *Fusarium graminearum*, vesicular trafficking, VIGS, wheat.

Introduction

In nature, plants are frequently exposed to various environmental stresses including pathogens, and yet more often than not plants appear healthy or show only weak or mild disease symptoms. To maintain this healthy status, plants have evolved an elaborate and tightly regulated innate immune system that allows them to restrict pathogen invasion or slow down/minimize disease progression (Jones and Dangl, 2006). These defensive processes not only include secretion of various peptides...
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and secondary metabolites in response to pathogen attack (Dixon, 2001; van Loon et al., 2006; Yun et al., 2013), but also the tight regulation of this secretion (Yun et al., 2013).

In addition to the cell wall, each plant cell is enclosed by the plasma membrane, and the cytoplasmic contents include a variety of membrane-enclosed organelles. Transport of various cargo molecules across different membranes and the sorting of these into the correct cellular compartments is a process central to the functioning of multiple plant cell types (Bassham et al., 2008). The transport of components in small, membrane-bound vesicles between the intra- and extracellular space is referred to as vesicular trafficking (Yun and Kwon, 2017). Regulation of multiple cellular responses by the membrane trafficking network during plant–microbe interactions is required to facilitate a coordinated defense response at sites of pathogen attack (Ben Khaled et al., 2015; Gu et al., 2017; Yun and Kwon, 2017; Ekanayake et al., 2019).

The vesicular trafficking system comprises two main pathways, secretory and endocytic (Fig. 1A), with both implicated in effective immunity against pathogens. A wide range of defense-related proteins, antimicrobial metabolites, and compounds such as callose that strengthen the plant cell wall can potentially be secreted at the sites of pathogen invasion (Gu et al., 2017). Concomitantly cell-surface immune receptors are subjected to endocytosis, which is necessary for initiation of signal transduction and regulation of receptor activity (e.g. through recycling or degradation in the vacuole) (Ranf, 2017). Secretery pathways transport newly synthesized proteins and other macromolecules (collectively referred to as ‘cargo’) from the endoplasmic reticulum via the Golgi apparatus to the plasma membrane or the extracellular space (Bassham et al.,...
Table 1. Arabidopsis mutants used in this study.

| Gene name       | Gene ID    | Gene description                                                                 | Mutant name | Mutant accession No. | Reference                          | Evidence in Arabidopsis plants                                                                 |
|-----------------|------------|----------------------------------------------------------------------------------|-------------|----------------------|------------------------------------|------------------------------------------------------------------------------------------------|
| Ara1/AtRabA5E   | At1g05810  | Arabidopsis gene homologous to the Ras oncogene 1/Rab GTPase-like A5A             | ara1        | SM_3.15870           | Karim et al. (2014)                 | Interacts with proteins involved in multiple stress responses                                 |
| Drp2b           | At1g59610  | Dynamin-related protein 2B                                                        | drp2b-1     | SALK_003049          | Smith et al. (2014)                 | Functions as a novel component of flagellin-mediated defense responses against bacterium *P. syringae* |
| Pen1/Syp121     | At3g11820  | Penetration resistance 1/Syntaxin of plants 121                                  | pen1        | SALK_004484C         | Collins et al. (2003), Johansson et al. (2014) | Absence of Pen1 is associated with increased tissue penetration by the non-adapted fungal pathogen *Blumeria graminis f.sp. hordei* |
| Syp122          | At3g52400  | Syntaxin of plants 122                                                            | syp122-1    | SALK_008617          | Zhang et al. (2007)                 | Acts together with SYP121 as a negative regulator of programmed cell death, salicylic acid, jasmonic acid, and ethylene pathways |
| AtMin7/Ben1/Big5| At3g43300  | HopM1 interactor 7/BFA-visualized endocytic trafficking defective 1/Brefeldin A-inhibited guanine nucleotide-exchange protein 5 | atmin7-1    | SALK_012013          | Nömura et al. (2006)                | Encodes an immunity-associated ARF-GEF family protein targeted by HopM1, conserved bacterial *P. syringae pv. tomato* virulence protein |
| Vps37           | At3g53120  | Vacuolar protein sorting 37                                                       | vps37-1     | SALK_042859C         | Spallek et al. (2013)               | Co-localizes with FLS2 (FLAGELLIN SENSING 2) immune receptor at endosomes and immuno-precipitates with this receptor upon flg22 elicitation |
| Vps28           | At4g05000  | Vacuolar protein sorting 28                                                       | vps28-2     | SALK_040274          | Spallek et al. (2013)               | Critical for immunity against bacterial infection through a role in stomatal closure; regulates late FLS2 endosomal sorting |
| Vamp721         | At1g04750  | Vesicle-associated membrane protein 721                                           | vamp721     | SALK_037273C         | Kwon et al. (2008b), Kim et al. (2014) | Forms a complex with PEN1; mediates trafficking of the powdery mildew resistance protein RPW8.2 to the extrahaustorial membrane of haustorial complexes |
| Lip5            | At4g26750  | LYST-interacting protein 5                                                        | lip5-1      | SAIL_854_F08         | Wang et al. (2014)                  | Disruption of Lip5 causes increased susceptibility to the bacterial pathogen *P. syringae* |
2008). In the endocytic pathway, membranous vesicles internalized in the plasma membrane undergo homotypic fusion to form early/sorting endosomes. Internalized cargo can then be sorted and recycled back to the plasma membrane through the recycling endosome, sent to the trans-Golgi network (TGN) via retrograde trafficking mechanisms, or trafficked through the late endosome/multivesicular body to the vacuole (Fig. 1A) (Kwon et al., 2008a, b; Dodds and Rathjen, 2010; Bozkurt et al., 2011; Nomura et al., 2011; Ellinger et al., 2013).

The increasing number of mutants in plant vesicular trafficking genes that show altered resistance to microbial pathogens illustrates the importance of vesicular trafficking to plant immunity (Rybak and Silke, 2019). Whereas the general understanding of these trafficking pathways has advanced, knowledge of the molecular mechanisms employed by plant cells to coordinate the transport of specific molecules through different cell compartments is still limited (Ekanayake et al., 2019). The components of vesicular trafficking do not function in isolation but form complexes. Within these complexes, a defined set of proteins are responsible for selecting and transporting specific cargo molecules in a way that is highly coordinated both temporally and spatially (Fig. 1B, C) (Rutter and Innes, 2018).

Plant pathogens are known to secrete small effector proteins some of which could specifically target components of vesicular trafficking important for plant immunity. One of these targets is the regulator protein adenosine diphosphate ribosylation factor–guanine nucleotide exchange factor (ARF-GEF) encoded by AtMin7/Big5 in Arabidopsis. The bacterial effector HopM1, which is required for proliferation of Pseudomonas syringae pv. tomato in the apoplast of the leaf interior, has been shown to degrade MIN7 in Arabidopsis. The bacterial effector HopM1, which is required for proliferation of P. syringae, ARF-GEF proteins are involved in controlling vesicle formation by regulating the ARF-family of small GTPases (Singh and Jürgens, 2018). Another example is the Phytophthora infestans effector AvR3a, which perturbs pattern-triggered immunity responses partly by targeting Dynamin-Related Protein 2B (DRP2B) in Arabidopsis (Chaparro-Garcia et al., 2015). Dynamins are important regulators of clathrin-mediated endocytosis and are involved in the scission and release of clathrin-coated vesicles from the plasma membrane (Gu et al., 2017).

Much of the evidence regarding the role of vesicular trafficking in plant immunity comes from studies involving the model plants Arabidopsis and Nicotiana benthamiana and to a lesser extent crop species such as barley (Hordeum vulgare), and a small number of their respective adapted biotrophic or hemi-biotrophic bacterial (P. syringae), oomycete (P. infestans) or ascomycete powdery mildew pathogens (Golovinomyces cichoracearum, Blumeria graminis, and Erysiphe pisi) (Nomura et al., 2006; An et al., 2006; Wang et al., 2009; Schmidt et al., 2014; Chaparro-Garcia et al., 2015). However, little is still known about the role of plant vesicular trafficking in the interactions involving other pathogens, other crop species, and those that primarily infect non-leaf tissue.

Fusarium head blight (FHB) disease, caused by the ascomycete fungus Fusarium graminearum and related Fusarium species, causes substantial yield losses and reduced grain quality and safety in a number of major cereal crops, such as wheat, barley, maize, rye, triticale, and oat, worldwide (McMullen et al., 2012; Dean et al., 2012). Moreover, under laboratory conditions F. graminearum is able to infect floral tissue of intact Arabidopsis plants as well as detached leaves (Chen et al., 2006; Cuzick et al., 2008; Brewer and Hammond-Kosack, 2015; Wood et al., 2020). For this model plant species, a large mutant collection is readily available.

The aim of this study was to assess whether knock-out mutations of the individual components of the vesicular trafficking system in Arabidopsis previously implicated in plant immunity had any impact on the interaction with F. graminearum. Screening of the assembled collection of mutants using a detached-leaf bioassay identified two independent T-DNA insertion mutants in the AtMin7 gene, which displayed striking hypersusceptibility to F. graminearum strain PH-1 compared with the parental wild-type Arabidopsis ecotype Col-0. Utilizing a recently released high-quality fully annotated wheat genome reference sequence assembly (International Wheat Genome Sequencing Consortium, 2018) and well-established bioinformatics tools enabling identification of putative gene orthologs from different plant species (Adamski et al., 2020), we identified the three homoeologous TaMin7 genes in hexaploid wheat (Triticum aestivum). Knock-down of these genes using virus-induced gene silencing (VIGS) (Lee et al., 2012) significantly promoted FHB disease formation in this crop species.

Materials and methods

Plant material and growth conditions

The Arabidopsis mutants (Table 1) and the corresponding wild-type parental ecotype Col-0 used in the study were obtained from the Nottingham Arabidopsis Stock Centre. The corresponding mutations were verified by PCR using the primers listed in Supplementary Table S1 and confirmed homozygous mutants selected for further study. Arabidopsis seeds were sown in Levington F2+S compost (Everris Ltd) and stratified in the dark for 4 d at 5°C before transferring to a controlled environment growth chamber operating at 20°C/17°C during day/night with a 12-h photoperiod (light intensity of approximately 80–100 µmol m−2 s−1).

The fully susceptible spring type bread wheat (Triticum aestivum) cv. Bobwhite was used in this study (Cuzick et al., 2008). The plants were grown in a controlled-environment growth chamber with day/night temperatures of 23°C/20°C at around 65% relative humidity and a 16-h photoperiod with light intensity of approximately 180 µmol m−2 s−1.

Fungal growth conditions, plants inoculation, and disease assessment

The reference F. graminearum strain PH-1 was used in this study. Routine culturing of the fungus, conidiospore induction, and preparation of conidial suspensions followed essentially the same procedures as described in Brown et al. (2011). Conidiospore suspensions harvested in
sterile distilled water were adjusted to a concentration of 5×10⁵ or 1×10⁶ conidia ml⁻¹ for inoculation of Arabidopsis or wheat, respectively.

Detached Arabidopsis leaves were inoculated as described in Chen et al. (2006) with the following modification. Fully expanded rosette leaves were detached from the 6-week-old plants using razor blades and placed adaxial surface facing upwards on the surface of 1% water agar in 10×10 cm square sterile Petri dishes, with eight leaves per dish. Each leaf was then superficially wounded by gently puncturing over the midrib with a glass Pasteur pipette and a 5 μl droplet of F. graminearum conidiospore suspension supplemented with 20 μM deoxyribonuclease (DON) was deposited onto the fresh wound. Mock inoculation was carried out using a 5 μl droplet of sterile distilled water supplemented with 20 μM DON. After inoculation, the plates were transferred to the controlled-environment growth chamber operating at 20°C/17°C during day/night and 16-h photoperiod but kept in the dark for the first 3 d following which plates were incubated under low light (40 μmol m⁻² s⁻¹) for a further 4 d before the disease assessment took place.

Color (RGB) photographs were taken at 6 d after inoculation using a Nikon (D90) camera and backlitting to ensure consistent illumination. Image analysis to quantify the diseased areas was conducted using the LemnaTec LennaGrid software module (LemnaTec GmbH, Aachen, Germany). Leaf areas were segmented using a combination of a color-based classification and thresholding after converting the images to grayscale. Filters were applied to remove misclassified pixels and to fill in gaps. Finally, a customized color-based classification was applied to score leaf-area pixels as belonging to diseased or healthy tissue. Intact spikes of adult wheat cv. Bobwhite plants were point inoculated at the first signs of anther extrusion by depositing 5 μl of conidial suspension in the floral cavity between the palea and lemma of the outer two florets located in the upper one-third of the spike, as previously described (Brown et al., 2011). Control plants were inoculated with sterile water only. Inoculated plants were incubated in a humid chamber for 48 h of which the first 24 h were in darkness. The inoculated plants were then kept in a controlled-environment growth chamber at approximately 65% humidity, the progression of the disease was visually monitored every 3 d, and the number of bleached spikelets below the inoculated spikelet on each spike was recorded along with the total number of spikelets on each spike (Urban et al., 2003).

PCR-based confirmation of specific mutations in the T-DNA insertion mutant Arabidopsis plants obtained from the seed stock center

One leaf from each Arabidopsis plant was collected in a 2 ml microtube, frozen in liquid nitrogen, and then ground with a micro-pestle. DNA extraction was carried as described in Motteram et al. (2009). Briefly, ground tissue was added to 350 μl of TEN buffer (500 mM NaCl, 400 mM Tris-HCl, 50 mM EDTA, pH 8.0), 1% β-mercaptoethanol, 5 mM 1,10-phenanthroline, and 2% (w/v) polyvinylpyrrolidone (K30). The resulting suspension was then mixed thoroughly with 350 μl of 2% (w/v) SDS and incubated for 30 min at 65°C. Following the addition of 300 μl of ice-cold ammonium acetate (7.5 M), the sample was kept on ice for 20 min and then centrifuged at 15 000 g for 10 min. DNA was precipitated with isopropanol, washed with 70% (v/v) ethanol, and dissolved in sterile distilled water.

For genotyping an insertion line using three primers in two combinations, we carried out two PCR reactions: the ‘wild-type PCR’ and the ‘T-DNA PCR’ (O’Malley et al., 2015). Primer sequences were available at the Salk Institute Genomic Analysis Laboratory (SIGnAL) database and are listed in Supplementary Table S1. The wild-type PCR reaction used gene-specific ‘right primer’ (RP) and ‘left primer’ (LP) that flank the T-DNA insertion site in the corresponding mutant and allowed amplification of DNA fragments from both wild-type plants and heterozygous mutants. In the T-DNA PCR reaction, a T-DNA left border (LB) primer and a gene-specific RP were used. This second PCR reaction selectively amplified the T-DNA–genomic DNA junction sequence and allowed amplification of DNA fragments from heterozygous and homozygous mutants. PCRs were performed using REDTaq ReadyMix PCR Reaction Mix (Sigma-Aldrich, Gillingham, UK) following the manufacturer’s instructions. A 1 μl aliquot of gDNA was used in a 25 μl PCR reaction, with an annealing temperature of 60°C. Primers were added at a final concentration of 0.5 μM.

Identification of wheat Min7 genes

Domain analysis of predicted ARF–GEF proteins in the hexaploid wheat (T. aestivum) was carried out using the BioMart tool in Ensembl. Initially searches were carried out for proteins that contained the Sec7_N domain (guanine nucleotide exchange factor in Golgi transport N-terminal domain; PF12783). The wheat genome assembly IWGSC RefSeq 1.0 (International Wheat Genome Sequencing Consortium, 2018) was used in this analysis. Coding sequences of eight previously identified genes comprising a small family of ARF–GEF encoding genes in Arabidopsis (Vernoud et al., 2003) were also extracted using the BioMart tool in Ensembl. Multiple protein sequence alignment was carried out using ClustalW tool in Geneious v.10. For phylogenetic reconstruction, the TVM+I+G nucleotide substitution model was selected by Akaike information criterion in jModeltest v.2.1.10 (Posada, 2008; Darriba et al., 2012). The maximum likelihood phylogeny was reconstructed using PhyML (Guindon and Gascuel, 2003), with the substitution model selected in jModeltest; starting tree with optimized topology, length, and rate parameters; topology searching by the best of nearest neighbor interchange and subtree pruning and regrafting; and 500 bootstraps.

Vector construction for virus-induced gene silencing

Total RNA extracted from healthy wheat cv. Bobwhite leaf tissue was used as a template for an RT-PCR amplification of a 209-bp TaMin7 gene fragment using primers TaMin7-2A-segl-R (5′-AACCACCACCC GTAAAGGTCGCCTCGTCAAT-3′) and TaMin7-2A-segl-F (5′-A AGGAAATTTAATGTGGCAAGCAAGCGCATC-3′). This fragment was cloned into an antisense orientation into the Barley stripe mosaic virus (BSMV) vector pCa-γbLIC using ligation–independent cloning (Yuan et al., 2011). The VIGS vector for silencing the control TaChH gene was kindly provided by Prof. Dawei Li (China Agricultural University, Beijing, China). BSMV:msc4D, containing a 275-nt non-coding DNA sequence), was used as a negative control (Saintenan et al., 2018). To prepare the virus inoculum for wheat inoculation, the BSMVβ, BSMVβ, and recombinant BSMVγ derivates containing msc4D or fragments of the TaMin7 or TaChH genes were transformed into the Agrobacterium tumefaciens strain GV3101 (pMP90). Agroinfiltration of Nicotiana benthamiana leaves was carried out as previously described (Lee et al., 2015). The agroinfiltrated N. benthamiana leaves were harvested 5–7 d post-agroinfiltration, homogenized in 10 mM Na–phosphate buffer (pH 6.8) containing 1% Celite 545 AW (Sigma–Aldrich), and the sap was mechanically inoculated onto wheat leaves just prior to appearance of a flag leaf. Fungal inoculation of wheat spikes at anthesis was then carried out as described above.

Quantification of gene expression and fungal biomass by quantitative real time PCR

For quantification of gene expression, total RNA was isolated from either Arabidopsis leaves or spike tissue of wheat plants using the TRIzol reagent (Thermo Fisher Scientific, Waltham, MA, USA) following the manufacturer’s instructions. To remove any traces of gDNA contamination, RNA samples were treated with TURBO DNaseI (Thermo Fisher Scientific) using methods as described by the manufacturer. The first
strand cDNA was synthesized from 1 µg of total RNA in a total volume of 20 µl using the SuperScript IV Reverse Transcriptase (Thermo Fisher Scientific) and oligo (dT)₁₈ primers according to the manufacturer’s instructions.

Gene-specific primers were used for RT-quantitative real time PCR (qPCR) analysis of transcript levels (Supplementary Table S1). Relative transcript levels were calculated by comparative threshold cycle (ΔΔCt) and normalized to the Arabidopsis ACTIN2 gene or the wheat CDC48 gene (Lee et al., 2014; Masachis et al., 2016).

A no template control was included in each of the qPCR experiments. For quantification of fungal biomass, total genomic DNA was extracted from infected leaves at 6 d post-inoculation using DNAeasy Plant Mini Kit (Qiagen, Manchester, UK) and subjected to qPCR using the primers specific for the F. graminearum ACTIN gene (Supplementary Table S1). Relative amounts of fungal DNA were calculated by comparative ΔΔCt and normalized to the Arabidopsis ACTIN2 gene (Masachis et al., 2016).

Protein extraction and western blots

Total leaf protein preparations were made as previously described (Sainsbury et al., 2009). Samples were resolved on 8% SDS-PAGE gels (Mini-PROTEAN, Bio-Rad) and transferred on to a nitrocellulose membrane (Hybond ECL, GE Healthcare). Immunoblots were performed by standard procedures using the Arabidopsis MIN7 specific antibodies at a dilution of 1:3000 (Nomura et al., 2006). The blots were developed using ECL Plus Western Blotting Detection Kit and images were acquired using Odyssey Imaging System (LI-COR Biosciences Ltd, Cambridge, UK).

Experimental design and statistical analysis

For the following experiments, GenStat for Windows (19th Edition, 2017; VSN International) was used. For Arabidopsis leaf inoculation assays, disease was quantified by expressing the diseased leaf area relative to the total leaf area. The statistical design for Arabidopsis leaf inoculation assays consisted of randomized blocks. Twelve leaves, one from each genotype, were placed onto each of eight plates. Seventy-two plants were used in total for each experiment (six plants per genotype). Six plates were used for fungal inoculation, and two plates were inoculated with water supplemented with DON to be used as a control. Three independent experiments were performed. Disease was quantified by expressing the diseased leaf area relative to the total leaf area. Mean disease levels for each genotype were compared using a multi-stratum ANOVA. Independent mutant genotypes were compared with the wild-type plants using a Dunnett’s test at the 5% (P < 0.05) level of significance using wild-type Arabidopsis Col-0 as the control test. GenStat (release 20.1, 2019; VSN International) was used for the statistical analyses. AtMin7 expression and fungal biomass determined by qPCR were compared from three biological replicates using ANOVA. Significance of differences between calculated means was determined using least significant difference (LSD) at the 5% level of significance.

For the statistical analysis of F. graminearum disease data, ANOVA was performed on the mean of the infected spikelets below the inoculation point out of the total spikelets calculated from the control (no virus and BSMV:ns-4D treated) and TdMIN7 silenced plants of three independent experiments, and linear models were fitted using GenStat. Graphical representations were made using the ggplot2 (Wickham, 2016) package in R.

Results

Assembling a collection of Arabidopsis mutants with defects in membrane trafficking

To investigate whether vesicular trafficking plays a role in a compatible interaction (i.e. disease formation) between the ascomycete fungus F. graminearum and its laboratory host Arabidopsis, we assembled a collection of 11 mutants containing T-DNA insertions in nine immunity-associated genes regulating different vesicular trafficking pathways (Fig. 1A; Table 1). Homozygous mutants were obtained from the Nottingham Arabidopsis Stock Centre (NASC, UK), and each mutant was verified by PCR amplification using gene-specific and T-DNA-specific primers (Supplementary Table S1).

Smaller rosettes were observed for three mutants (vump721, atmin7-1, and atmin7-2) when compared with the corresponding wild-type Arabidopsis ecotype Col-0 plants grown under standard controlled-environment conditions. For the remaining mutants, no obvious developmental or growth defects were observed. Representative images of each mutant compared with Col-0 at two different growth stages are given in Supplementary Fig. S1.

MIN7, an ARF-GEF protein, is required to minimize F. graminearum infections in Arabidopsis

To gain insight into whether mutations in any of the selected vesicular trafficking genes increase or decrease susceptibility to the virulent F. graminearum strain PH-1, young 6-week-old Arabidopsis plants were point inoculated with F. graminearum conidial suspension supplemented with the mycotoxin DON in a detached leaf bioassay (Chen et al., 2006). At 6 d post-inoculation (dpi), the inoculated leaves were photographed, and disease levels were quantified by measuring the proportion of lesioned/necrotic area compared with the total leaf area by analysing the images using the LennaGrid software module (LemnaTec GmbH, Aachen, Germany).

Fusarium graminearum–inoculated leaves of the two independent T-DNA insertion mutants in the AtMin7 gene (At3g43300) developed extensive necrotic lesions covering up to 100% of the total leaf area and showed almost complete loss of green photosynthetic tissue, while most of the remaining mutants displayed much milder disease symptoms with smaller lesions (Fig. 2A, B). Quantification of F. graminearum biomass by qPCR showed a markedly higher fungal burden in atmin7-1 and atmin7-2 mutants compared with wild-type Col-0 or any of the other analysed mutants at 6 dpi (Fig. 2C). These results strongly suggest that plants with knock-out mutations in the AtMin7 gene are significantly more susceptible to F. graminearum.

AtMin7 is a large gene of 5857 bp containing 33 exons, and is located on chromosome 3. The two loss of function mutants studied here, atmin7-1 and atmin7-2, carry T-DNA insertions in exon 1 and exon 18, respectively (Nomura et al., 2006). AtMin7 is a member of a small family comprising eight genes encoding ARF-GEF proteins (Supplementary Fig. S2), which play important roles in the budding of transport vesicles from the membranes (Fig. 1A) (Steinmann et al., 1999; Mosessova et al., 2003). This vesicular trafficking component has been shown to contribute to resistance to the bacterial pathogen P. syringae pv.
tomato, possibly through regulating the trafficking of immunity-associated cargo molecules (Nomura et al., 2006, 2011).

**Expression analysis of AtMin7 in Arabidopsis leaves infected with F. graminearum**

To understand further the mechanisms by which disruption of AtMin7 led to Arabidopsis susceptibility to F. graminearum, expression of AtMin7 was measured in wild-type Col-0 leaves inoculated with fungal spores. Transcripts levels were compared with mock-inoculated leaves (water and DON). Although considerable variation in AtMin7 expression was noted from the fungal inoculated leaves between replicates, no significant differences were observed between the mock and F. graminearum treatments (Fig. 3A). Therefore, F. graminearum infection does not seem to suppress expression of AtMin7 gene and/or transcript abundance during the infection cycle on Arabidopsis leaves.

![Fig. 2. Fusarium graminearum infection of different Arabidopsis mutants. (A) Box-plot and dot-plot of Arabidopsis detached leaves infected with F. graminearum. Disease was measured as the proportion of infected leaf area compared with the total leaf area. Symbols indicate treatments statistically different from Col-0 (WT) control according to Dunnett’s test ($P < 0.05$). Data represent results from three different experiments with at least 72 leaves from 12 different plants from each mutant for each experiment. (B) The appearance of representative detached leaves 6 d after spore droplet inoculation. (C) Representative photos of leaves mock and F. graminearum inoculated. Mock treatments were with water supplemented with 20 µM DON. (D) Fungal biomass determined by qPCR. The relative amount of fungal DNA was calculated using the threshold cycle ($\Delta\Delta C_t$) method, normalized to the Arabidopsis ACTIN2 gene ($P < 0.05$; mutant versus wild-type according to the least significant difference, LSD). $n = 3$ biological replicates.](image)
We next asked if MIN7 protein levels would also remain unchanged in *F. graminearum*-inoculated leaves. To test this, MIN7-specific antibodies were used to detect the protein in Arabidopsis leaves inoculated with *F. graminearum*, mock inoculated or left untreated (Fig. 3). MIN7 was detected in untreated and mock-inoculated leaves, whereas this protein was practically undetectable in the *F. graminearum*-inoculated leaves (Fig. 3B). These results suggest that MIN7 is destabilized and/or degraded during fungal infection.

**Identification and expression analysis of candidate wheat TaMin7 genes**

To identify homologs of *AtMin7* gene in wheat, a natural crop host of *F. graminearum*, we used the BioMart data mining tool available through Ensembl Plants (Smedley et al., 2015). A total of 26 gene sequences were identified in the reference hexaploid wheat (*Triticum aestivum*) genome (*2n* = 6x = 42, AABBDD) by searching for genes encoding proteins containing the catalytic SEC7 domain (PF12783) characteristic of ARF-GEF proteins. We then aligned the proteins encoded by the identified wheat genes with all eight members of the Arabidopsis ARF-GEF family proteins and used the resulting multiple alignment for phylogenetic analysis. The constructed maximum likelihood phylogenetic tree revealed that the three closely sequence-related wheat proteins formed a distinct clade with Arabidopsis MIN7, suggesting these proteins represent the wheat A-, B- and D-genome-encoded orthologs of MIN7 (Fig. 4).

TaMin7 transcript levels were also measured in wheat spikes both inoculated and non-inoculated with *F. graminearum*. Wheat spikelets were point inoculated with either water or fungal spores and tissues were collected at 5 dpi for RNA extraction and gene expression analysis using qPCR. Similar to Arabidopsis *AtMin7*, no significant differences were observed in the expression of *TaMin7* genes between mock and fungal inoculated wheat plants (Supplementary Fig. S3). Unfortunately, the lack of antibody specificity for wheat MIN7 prevented protein quantification assessments in wheat.

**Silencing the candidate TaMin7 genes in wheat spikes enhances susceptibility to *F. graminearum***

To explore the potential function for MIN7 in the *F. graminearum*-wheat interaction, we tested the effect of silencing the three homoeologous *TaMin7* genes (*TraesCS2A02G202900, TraesCS2B02G230000, and TraesCS2D02G212800*) using BSMV-mediated VIGS on FHB disease development (Lee et al., 2012). A 209 bp region highly conserved between coding sequences of the three *TaMin7* homoeologs was selected as a target for VIGS using the si-Fi21 software (Lück et al., 2019). This target region was predicted to generate a high number of silencing-effective siRNAs (*n* = 91), and a very low likelihood of off-target silencing.

An important factor for successful application of VIGS is the ability of the virus to infect and spread without having any deleterious effect in the host plant. Therefore, the feasibility of using the BSMV-mediated VIGS approach to induce systemic silencing in the spike tissue of wheat cv. Bobwhite susceptible to *F. graminearum* PH-1 (Brown et al., 2011) was first tested by visualizing the phenotype induced by silencing the *Magnesium-chelatase subunit H* (*TaChlH; TraesCS7A02G480700*) gene. *TaChlH* is involved in chlorophyll biosynthesis and is often...
used as a visible marker in VIGS (Yuan et al., 2011). The recombinant BSMV carrying a 250 bp fragment of TaChlH gene in an antisense orientation and BSMV::mcs4D harboring a non-coding DNA sequence of 275 bp were inoculated onto the flag leaves of wheat plants at the early boot stage. At 13 dpi the plants infected with BSMV::asTaChlH developed yellow-orange coloration of the lemma and palea of spikes indicative of the loss of chlorophyll and successful silencing of the TaChlH gene, whereas the spikes of plants infected with the control construct BSMV::mcs4D showed typical mosaic symptoms (Supplementary Fig. S4). No visible developmental abnormalities were observed in the wheat plants challenged with either of the two VIGS constructs. Moreover, similar levels of FHB disease were observed on wheat plants pre-infected with BSMV and on virus-free plants challenged with F. graminearum, indicating that the susceptibility to the fungus was not compromised in the virus-infected wheat (Fig. 5). These results suggest that BSMV-mediated VIGS can be used to silence genes in the reproductive tissue of wheat cv. Bobwhite and the approach appeared suitable for assessing the role of TaMin7 during F. graminearum infection in wheat.

The ability of BSMV::asTaMin7 to induce silencing of the corresponding endogenous genes was confirmed using RT-qPCR. The TaMin7 transcripts were decreased in abundance by 77% in the spikes of the VIGS-treated plants compared with those inoculated with the negative control BSMV::mcs4D, and no significant difference in TaMin7 expression levels was observed in the mock- versus BSMV::mcs4D-inoculated plants. To analyse the effect of TaMin7 silencing on F. graminearum infection, spikes of VIGS-treated plants showing typical virus-induced symptoms were point inoculated with the fungal conidia suspension and observed for disease symptoms for 15 d after fungal inoculation. Reduction of TaMin7 mRNA expression levels in spikes of the BSMV::asTaMin7-treated plants was associated with significantly enhanced susceptibility to F. graminearum (Fig. 5). In contrast, no effect on FHB disease development was found in mock-inoculated and BSMV::mcs4D-inoculated control plants (Fig. 5).

**Discussion**

In this study, we have compared disease symptom formation caused by the fungus F. graminearum on the leaves of 11 different Arabidopsis vesicle trafficking mutant lines available in the ecotype Col-0 background. In total, eight different components of the membrane trafficking system, previously demonstrated to play important role(s) in plant defense during plant–pathogen interactions, have been evaluated (Table 1) (Gu et al., 2017; Yun and Kwon, 2017; Ekanayake et al., 2019). Strikingly, the two independent atmin7 mutants were shown to be highly susceptible to F. graminearum infection. Similarly, when expression of the three orthologs of AtMin7 were knocked-down in hexaploid wheat through VIGS, these TaMin7 silenced plants displayed significantly more FHB disease formation. Quantification of AtMin7 transcript levels in Arabidopsis and TaMin7 in wheat revealed no evidence that F. graminearum infections caused a reduction in transcript abundance. Instead, immunological evidence in Arabidopsis suggested that F. graminearum infections cause a reduction in MIN7 protein levels.

Some bacterial and oomycete pathogens are known to have evolved effector proteins that are translocated into plant cells to promote disease formation by interfering with plant membrane trafficking pathways (Ben Khaled et al., 2015). One of the vesicular trafficking genes whose role during plant disease has been previously well explored in the interactions...
New role for MIN7 in minimizing fungal infection

between Arabidopsis and the bacterial pathogen *P. syringae* pv. *tomato* is AtMIN7 (Nomura et al., 2006, 2011). The MIN7 protein localizes to the TGN/early endosome and is involved in endocytic recycling of plasma membrane resident proteins but it has also been hypothesized to regulate secretion (Tanaka et al., 2009; LaMontagne and Heese, 2017) (Fig. 1). Mutants that lack this protein allow increased bacterial multiplication, possibly due to misregulation of membrane trafficking of the plant defense-related cargo (Nomura et al., 2006, 2011). MIN7 has also been shown to contribute to the cytosol-initiated immune responses triggered by the *P. syringae* pv. *tomato* effectors such as AvrRpt2 and AvrPphB (Nomura et al., 2011) and to the apoplast-initiated immune responses through an unknown mechanism by preventing apoplast water soaking and therefore presumably restricting the flow of nutrients to the bacteria (Xin et al., 2016). To achieve successful disease...
lesion formation, *P. syringae pv. tomato* secretes a conserved effector protein, HopM1, which is translocated to the TGN/early endosome of its host during infection where it mediates destabilization of MIN7 followed by its degradation via the 26S proteasome (Nomura et al., 2011). Here we clearly demonstrated that MIN7 also contributes to defense against the fungal pathogen *F. graminearum* because the absence of this protein in Arabidopsis resulted in a markedly enhanced disease (Fig. 2). However, whereas atmin7 mutants displayed increased susceptibility to the *P. syringae pv. tomato* ∆cml mutant that lacks HopM1 along with several other conserved effectors including AvrE (Nomura et al., 2011), we show that these same Arabidopsis mutants are clearly and unmistakably hypersusceptible to the wild-type strain of *F. graminearum* (Fig. 2).

Disruption of AtMin7 may compromise trafficking of specific molecules and cargo protein. These molecules could include plant defense-related proteins and/or secondary metabolites that help minimize *F. graminearum* infection. For example, callose, a (1,3)-β-glucan polymer, can act as a physical barrier reducing fungal penetration. Both increased callose deposition and *F. graminearum*–induced callose synthase activity in the infected spikelets and rachis nodes were found to be correlated with increased disease resistance (Ribichich et al., 2000; Blümke et al., 2014). Identification of proteins and/or other molecules transported specifically by membrane vesicles regulated by MIN7 is challenging, but comparison of vesicles cargo between wild-type and atmin7 Arabidopsis mutants during *F. graminearum* infection could provide valuable clues to the role of MIN7 protein during fungal disease establishment.

Surprisingly, none of the other tested trafficking Arabidopsis mutants gave a distinct phenotype when detached leaves were challenged with *F. graminearum* spores (Fig. 2A). Most of the tested mutants, dpr2b-1, dpr2b-2, vps37-1, vps28-2, lip5-1, and AtMin7, were previously reported to be involved in immune responses to bacterial infection or bacterial pathogen–associated molecular patterns. DRP2B, VPS37, and VPS28 appear to be required for the regulation of the cell surface innate immune receptor FLAGELLIN SENSING 2 (Spellek et al., 2013; Smith et al., 2014). Disruption of LIPS compromises basal resistance to *P. syringae* (Wang et al., 2014). Therefore, it is possible that this sub-set of trafficking proteins in Arabidopsis may have specialized function(s) in response to bacterial infection, but not fungal invasion. An unexpected result was obtained with the pen1 mutant (Fig. 2A), which encodes the first SNARE to be identified with an immune function. PENETRATION-1 (PEN1), also known as Qa-SNARE SYNTAXIN OF PLANTS 121 (SYP121), is a component of a complex of SNARE proteins that plays a role in ‘point defense’ against fungal invaders, and loss of PEN1 function in Arabidopsis leads to almost 90% penetration success of the spores of the non-adapted powdery mildew fungus *Blumeria graminis* f. sp. *hordei* (Collins et al., 2003). PEN1/SYP121 forms a complex with at least four other proteins, namely SYNAPTOSOMAL-ASSOCIATED PROTEIN 33 kDa (SNAP33), Qbc–SNARE, and R–SNARE proteins VESICLE-ASSOCIATED MEMBRANE PROTEIN 721 and 722 (VAMP721 and VAMP722). We also tested sensitivity of vmp721 to *F. graminearum* in our study and observed levels of infection undistinguished from those in the wild-type Arabidopsis plants. However, VAMP721 and VAMP722 are known to be functionally redundant. The vmp721 vmp722 double null mutant plants could not be tested in the *F. graminearum* leaf bioassay because of severe growth defects and sometimes seedling lethality (Kwon et al., 2008; Yun et al., 2013). Finally, in this study of *F. graminearum*, the level of infection of the syp122 mutant was comparable to that in the wild-type Arabidopsis plants (Fig. 2A). SYP122 has been shown to share an overlapping function with PEN1 during certain aspects of plant development. However, when syp122 mutant plants were tested for resistance to *B. graminis* f. sp. *hordei*, a wild-type-like resistance response was retained (Assaad et al., 2004). Collectively, these results indicate that there are differences in the function of individual vesicular trafficking proteins when defending against adapted or non-adapted fungal pathogens.

Although Arabidopsis has proven to be a very useful model organism for unraveling the key mechanisms underlying interactions with pathogens including *F. graminearum* (Brewer and Hammond-Kosack, 2015), some findings cannot be translated directly to crop plants. Hence, studies involving interactions between the pathogens and their natural host plants could provide more relevant information and facilitate exploration of new strategies for disease control in crops. Here we utilized a transient gene silencing approach (VIGS) to assess the role of AtMin7 homologs in wheat, which is one of the most important staple food crops whose production is regularly threatened by fungal diseases including FHB. The results obtained from the analysis of the wheat–*F. graminearum* floral pathosystem (Fig. 5) are consistent with those from the study of the Arabidopsis–*F. graminearum* leaf pathosystem (Fig. 2) and provide evidence that disruption of MIN7 function in both dicotyledonous and monocotyledonous hosts and in both floral and leaf tissues compromises the plant innate immunity resulting in more severe disease.

To investigate whether *F. graminearum* utilizes similar mechanisms to the bacterial pathogen *P. syringae pv. tomato* to suppress host immunity and promote disease formation, the abundance of the *AtMin7* transcripts and MIN7 protein was determined (Fig. 3; Supplementary Fig. S3). In the various published whole wheat spike–*F. graminearum* transcriptome analyses, the abundance of the *TaMin7* transcripts was found to be highly variable. For example, transcriptome analysis revealed less than a 1 log2-fold change of *TaMin7* in both inoculated and non-inoculated plants in fully susceptible, moderately susceptible, or resistant cultivars (Pan et al., 2018). The RT-qPCR analysis done in this study confirms that *TaMin7* transcript abundance is roughly equivalent in inoculated and non-inoculated whole wheat spikes collected at
5 dpi (Supplementary Fig. S3). However, in a previously published transcriptome study that focused on dissected out different phases of *F. graminearum* infection, TaMin7 was down-regulated in infected tissue but only during the later symptomatic phase of disease formation (Dilks et al., 2019). In contrast, western blot analysis with anti-MIN7 antibodies provided clear evidence that MIN7 protein levels decrease following *F. graminearum* infection of Arabidopsis leaves (Fig. S3). Therefore, it is tempting to speculate that *F. graminearum* may contain effector(s) functionally similar to the *P. syringae pv. tomato* effector HopM1 that destabilizes MIN7 to suppress host immunity and hence to promote disease. *Fusarium graminearum* does not appear to possess a homolog of HopM1, and therefore further studies would be necessary to prove or disprove the above hypothesis.

Arabidopsis MIN7 has been shown to play a role in polar localization and dynamic repolarization of the PIN (PIN-formed) efflux carrier proteins enabling the directional transport of auxin in the tissues (Tanaka et al., 2013). Elevated levels of reactive oxygen species induced during stress responses in Arabidopsis affect MIN7-dependent PIN endocytic recycling, resulting in increased accumulation of auxin in the affected tissues (Zwiewka et al., 2019). A recent study demonstrated that higher levels of auxin are accumulated during *F. graminearum* infection in a susceptible wheat cv. Roblin compared with the moderately resistant cultivars Wuhan 1 and Nyubai, indicating that auxin may promote susceptibility to this fungal pathogen (Brauer et al., 2019). However, prior exogenous application of auxin increased resistance to both floral and root *Fusarium* diseases whilst cytokinin applications increased both tissues’ susceptibility (Petti et al., 2012, Haidoulis and Nicholson, 2020). Collectively, the data from these previous studies together with findings from this study form the foundations of an alternative hypothesis regarding the mechanisms employed by *F. graminearum* for achieving successful infection. That is, it is conceivable that *F. graminearum* upon infection induces temporally coordinated waves of gene expression that regulate MIN7-dependent distribution and accumulation of auxin during infection, whereas the artificial elevation of auxin levels prior to infection significantly disrupts the establishment of this subtle *F. graminearum* controlled hormonal changes. Further studies are necessary to confirm or refute this hypothesis.

In this study, the initial vesicular trafficking mutant screen was done using Arabidopsis leaf infections. This approach has many advantages but also some drawbacks. For example, could some of the interaction phenotypes and mechanisms specific to *F. graminearum* infection of wheat spike tissues have been missed by the experimental approach taken? In the last three years considerable advances have been made in the resources and technologies now available to the global wheat research community. For example, fully sequenced wheat genome and transcriptomes are available for the reference wheat genotype Chinese Spring as well as multiple wheat cultivars (International Wheat Genome Sequencing Consortium, 2018; Walkowiak et al., 2020). In addition, mutant targeting induced local lesions in genomes (TILLING) populations and genome editing technologies can now be used in addition to VIGS for functional analyses in both hexaploid and tetraploid wheat species (Krasileva et al., 2017). In the near future, by selecting and purifying the most appropriate lines from these new resources it should be possible to re-evaluate in both tetraploid and hexaploid wheat the various role(s) of the full spectrum of vesicular trafficking mutants against a range of diverse pathogens with different *in planta* lifestyles that also infect different host niches.

Elucidating the mechanisms by which membrane trafficking regulates plant immune responses and acquiring an enhanced understanding of the membrane trafficking components and pathways manipulated by microbial pathogens to promote disease will provide fundamental new knowledge for the development of novel methods of disease intervention.

### Supplementary data

The following supplementary data are available at *JXB* online.

Fig. S1. Representative photos of Arabidopsis mutants and wild-type Col-0 used in this study.

Fig. S2. Maximum likelihood phylogenetic tree indicating the relationship among Arabidopsis ARF-GEF encoding genes (coding sequences only).

Fig. S3. Expression analysis of *TaMin7* homologous genes in wheat spikes mock and *F. graminearum* inoculated by RT-qPCR.

Fig. S4. Silencing of *TaChlH (Mg-chelatase subunit H)* gene in wheat spikes.

Table S1. Primers used in this study for genotyping the various Arabidopsis mutants and for PCR analysis.

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### Author contributions

KK and KEH-K conceived the original idea and formulated a research plan; AKMW designed and performed most of the experiments and analysed the data; VP carried out the VIGS’ experiments in wheat and analysed the data; MGM provided technical assistance to AKMW; TA developed an image processing pipeline for quantifying fungal disease symptoms in Arabidopsis leaves; AKMW, VP, and KK wrote the manuscript with inputs from all the authors.
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