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**Phytophthora infestans** effector SFI3 targets potato UBK to suppress early immune transcriptional responses

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**Summary**

- The potato blight agent *Phytophthora infestans* secretes a range of RXLR effectors to promote disease. Recent evidence indicates that some effectors suppress early pattern-triggered immunity (PTI) following perception of microbe-associated molecular patterns (MAMPs). *Phytophthora infestans* effector PiSFI3/Pi06087/PexRD16 has been previously shown to suppress MAMP-triggered pFRK1-Luciferase reporter gene activity. How PiSFI3 suppresses immunity is unknown.
- We employed yeast-two-hybrid (Y2H) assays, co-immunoprecipitation, transcriptional silencing by RNA interference and virus-induced gene silencing (VIGS), and X-ray crystallography for structure-guided mutagenesis, to investigate the function of PiSFI3 in targeting a plant U-box-kinase protein (STUBK) to suppress immunity.
- We discovered that PiSFI3 is active in the host nucleus and interacts in yeast and in planta with STUBK. UBK is a positive regulator of specific PTI pathways in both potato and *Nicotiana benthamiana*. Importantly, it contributes to early transcriptional responses that are suppressed by PiSFI3. PiSFI3 forms an unusual trans-homodimer. Mutation to disrupt dimerization prevents nucleolar localisation of PiSFI3 and attenuates both its interaction with STUBK and its ability to enhance *P. infestans* leaf colonisation.
- PiSFI3 is a ‘WY-domain’ RXLR effector that forms a novel trans-homodimer which is required for its ability to suppress PTI via interaction with the U-box-kinase protein STUBK.

**Introduction**

The plant immune system may be activated following the detection of conserved microbial molecules (microbe-associated molecular patterns, MAMPs) by cell surface pattern recognition receptors (PRRs). Pathogen effectors serve to suppress this pattern-triggered immunity (PTI), or otherwise manipulate processes in the host to promote susceptibility. Effectors may be delivered by microbial pathogens to act in the apoplast or within plant cells (Jones & Dangl, 2006).

The late blight pathogen *Phytophthora infestans* remains a constant threat to potato and tomato production, precipitating considerable crop losses annually (Fry et al., 2015; Kamoun et al., 2015). Amongst a large repertoire of virulence determinants are the RXLR effectors, so-called because of the conserved Arg–any amino acid–Leu–Arg motif that facilitates their delivery inside plant cells (Rehmann et al., 2005; Whisson et al., 2007; Wawra et al., 2017). RXLR effectors manipulate a range of host processes by direct interaction with diverse plant proteins (Whisson et al., 2016). Some host proteins targeted by *P. infestans* RXLR effectors are positive regulators of immunity that confer posttranslational modifications (PTMs). For example, the RXLR effector AVR3a targets CMPG1, a host ubiquitin E3 ligase required for cell death triggered by perception of the PAMP infestin-1 (INF1; Bos et al., 2010) and a range of additional pathogen elicitors perceived at the cell surface (Gilroy et al., 2011). RXLR effector PexRD2 targets the kinase domain of the host protein MAP3Kɛ, which mediates signal transduction following perception of *Cladosporium fulvum* apoplastic effector Avr4 by the tomato Cf4 resistance protein (King et al., 2014).

By contrast to these examples, *P. infestans* RXLR effectors also target host proteins conferring PTMs that act as negative regulators of immunity, the so-called susceptibility (S) factors (Boevink et al., 2016a). For example, effector Pi02860 suppresses INF1-mediated cell death and interacts with host protein StNRL1, which is predicted to complex with Cullin 3 to act as an E3 ligase (Yang et al., 2016). However, StNRL1 is a negative regulator of INF1-mediated cell death, and therefore Pi02860 promotes its activity in targeting the positive immune regulator SWAP70 for proteasome-mediated degradation (He et al., 2018). In addition,
whereas PexRD2 suppresses the kinase signalling activity of its target MAP3K (King et al., 2014) to prevent phosphorylation of its substrates, RXLR effector Pi04314 interacts with protein phosphatase 1 (PP1) catalytic isoforms to form holoenzymes that are predicted to dephosphorylate substrates (Boevink et al., 2016b), and effector Pi17316 targets the MAP3K VIK to support its activity as a negative regulator of immunity (Murphy et al., 2018). These examples demonstrate that oomycetes can alter PTMs that positively or negatively regulate immunity.

Phosphorylation and ubiquitination are pervasive PTMs that affect all processes inside eukaryotic cells. The interplay and cross-talk between the two events have become a recurrent theme in cell signalling regulation (Nguyen et al., 2013; Filipčík et al., 2017). Protein kinases and ubiquitin E3 ligases play roles in both positive and negative regulation of PTI and ETI. Many PRRs involved in pathogen detection, for example, are kinases that initiate phosphorylation cascades (e.g. Wang et al., 2018). PRRs activate mitogen-activated protein kinase (MAPK) cascades involved in regulating immunity (Mithoe & Menke, 2018). In addition, kinases activate transcriptional regulators of immunity (Ishihama & Yoshioka, 2012), and regulate the complex crosstalk between phytohormone signalling pathways (Chen et al., 2010b). On the ubiquitination side, in addition to CMPG1 and NRL1, another E3 ubiquitin ligase, PUB17, functions in the host nucleus to mediate both PTI, following perception of the bacterial PAMP flg22, and cell death triggered by expression of Cf4/Avr4 but not INF1 (He et al., 2015). By contrast with CMPG1 and PUB17, some PUB E3 ligases have been shown to negatively regulate immunity. Two U-box E3 ligases, PUB12 and PUB13, negatively regulate flagellin (flg22)-triggered defense responses via the ubiquitination-mediated turnover of FLS2 (Lu et al., 2011; Kong et al., 2015). PUB22 targets a subunit of the exocyst complex, Exo70B2 for proteasomal degradation required for PAMP-triggered responses in Arabidopsis (Stegmann et al., 2012). More recently, the substrate adaptor component of a CULLIN3-based E3 ligase, POB1, has been shown to negatively regulate a range of immune responses and, indeed, one of its substrates for proteasome-mediated degradation is PUB17 (Orosa et al., 2017).

In this study, we focused on an RXLR effector from P. infestans, SF13/Pi06087/PexRD16, which is upregulated during the biotrophic phase of leaf and tuber infection in strains from diverse geographical locations (Haas et al., 2009; Oh et al., 2009; Cooke et al., 2012; Ah-Fong et al., 2017; Yin et al., 2017). Unlike PiSFI5, which suppresses flg22-triggered immunity upstream of MAP kinase activation by interacting with calmodulin (Zheng et al., 2018), PiSFI3 was able to suppress flg22-induced promoter-FLG22-INDUCED RECEPTOR-LIKE KINASE1 – LUCIFERASE (pFRK1-Luc) reporter gene activity downstream of MAP kinase activation in protoplasts of the host plant tomato (Zheng et al., 2014). Here, we show that when expressed in N. benthamiana or potato, PiSFI3 acts in the nucleus to enhance P. infestans leaf colonization. Overexpression of PiSFI3 in transgenic potato lines suppressed early transcriptional responses triggered by flg22. PiSFI3 interacts with a protein containing both U-box and kinase domains, StUBK, in yeast-two-hybrid (Y2H) assays and in planta. Silencing of UBK in N. benthamiana by virus-induced gene silencing (VIGS) or in potato by RNAi enhanced P. infestans leaf colonization. Transient expression of StUBK reduces P. infestans colonization and prevents PiSFI3 virulence activity. Moreover, VIGS or RNAi of UBK suppresses early transcriptional responses triggered by flg22 but has no effect on the cell death triggered by P. infestans PAMP INF1 and Cf4/Avr4 co-expression. The crystal structure of PiSFI3 confirms it possesses the WY-domain fold (Boutemy et al., 2011), but adopts an unusual trans-homodimer and can oligomerise in planta. Mutation of pairs of Leu/Asp residues that face each other across the PiSFI3 α-helices, disrupted oligomerisation and attenuated virulence function. We further found that PiSFI3 Leu/Asp mutants failed to localize to the nucleus and were not able to interact with StUBK, demonstrating that interaction with StUBK is required for the virulence function of PiSFI3.

Materials and Methods

Plasmid constructs

The full-length StUBK gene was cloned from potato cDNA with gene-specific primers modified to contain the Gateway® (Invitrogen) attB recombination sites. Polymerase chain reaction (PCR) products were purified and recombined into pDONR201 (Invitrogen) to generate entry clones. SF13, minus the signal peptide-encoding portion, was amplified from P. infestans cDNA and cloned into pDONR201. Primer sequences are shown in Supporting Information Table S1. Protein fusions were made by recombining the entry clones with the following plant expression vectors: pB7WGF2, pK7WGR2 and pGWB18 to generate N-terminal green fluorescent protein (GFP), monomeric red fluorescent protein (mRFP) and CMyc fusions respectively. SF13 double mutant clones were obtained by gene synthesis (Genesys, Genscript HK Ltd, Hong Kong, China) and cloned as above.

Potato transformation and plant growth condition

The RNAi construct was designed in the nonconserved 5′ regions of NlUBK and cloned into the expression vector pHell8 gate 8. Primer sequences are shown in Table S1. Agrobacterium tumefaciens containing the RNAi vector was transformed into the potato cultivar E3 by microtuber disc transformation as described by Si et al. (2003). Putative transgenic potato plants harbouring the target gene construct were screened on differential medium (3% Murashig and Skoog medium (MS) + 0.2 mg l−1 indole-3-acetic acid + 0.2 mg l−1 gibberellic acid A3 + 0.5 mg l−1 6-Benzylaminopurine + 2 mg l−1 zeatin + 75 mg l−1 kanamycin (kan) + 200 mg l−1 cephaplorins (Cef), pH 5.9) and then transferred to root generation medium (3% MS + 50 mg l−1 kan + 400 Cef mg l−1, pH 5.9). Transformants were confirmed by PCR with a forward primer binding to the 35S promoter and a gene-specific reverse primer. Gene expression levels were analyzed by qRT-PCR. The potato plantlets were propagated in MS medium (4% sucrose and 0.7% agar) and raised in a climate-controlled room at 20°C (16 h : 8 h, light : dark cycle).
**Phytophthora infestans** infection assay

*Phytophthora infestans* 88069dGT10 was grown in Petri dishes (90 mm diameter) of rye agar medium supplemented with geneticin antibiotic at 19°C. The *P. infestans* strain, Ljx18 (race 3.4.7.10.11) and HB09-14-2 (race 1.2.3.4.5.6.7.9.10.11, collected from Hubei Province, China), were cultured and propagated on rye agar at 19°C. Spores were prepared from cultures grown for 13 d. The spore concentration for the inoculum was adjusted by dilution in sterile distilled water to yield $4 \times 10^5$ sporangia ml$^{-1}$ for *N. benthamiana* leaves (VIGS leaves and transient expression assays), higher inoculum ($8 \times 10^4$ sporangia ml$^{-1}$) for GFP–StUBK transient expression assays and infection, and $7 \times 10^4$ sporangia ml$^{-1}$ for potato leaves. Next, 10 μl droplets were inoculated onto the abaxial side of detached *N. benthamiana* leaves (four sites per leaf) or potato leaves (two sites per leaf). Inoculated leaves were stored on moist tissue in sealed boxes. The infection assays were carried out as described previously (He et al., 2015; Boevink et al., 2016b; Yang et al., 2016).

**Tobacco rattle virus (TRV)-based VIGS in N. benthamiana**

Two portions of the nonconserved 5′ region of *NbUBK* were selected to make VIGS constructs. Regions were selected that shared no significant homology with other sequences in *N. benthamiana* to minimise the potential for off-target silencing. The two regions of *NbUBK* were amplified from *N. benthamiana* cDNA and cloned into pBinary TRV vectors (Liu et al., 2007) between *HpaI* and *EcoRI* restriction enzyme recognition sites in the antisense orientation. A TRV construct expressing GFP was selected to make VIGS constructs. Regions were selected that predicted extinction coefficients for the fusion and cleaved proteins of 19 940 M$^{-1}$ cm$^{-1}$.

**Heterologous expression in E. coli, protein production and purification**

SFI3 lacking the first 48 N-terminal amino acids (signal peptide and RXLR sequences removed), was cloned into the pOPINF plasmid (Berrow et al., 2007) by In-Fusion cloning (Clontech) following PCR using forward (AAGTTCTGTTTCAAGGG CCGTCTGATTTCGGCCAATTTCGCGCGGCGG) and reverse (ATGGTCTAGAAGCCTTAAGGTTGCGCGG CCGTCTGATTTCGGCCAATTTCGCGCGGCGG) primers (Integrated DNA Technologies).

SFI3 protein was produced in *E. coli* BL21 (DE3) cells grown in Luria-Bertani medium until optical density (OD) reached OD$_{600nm}$ 0.5–0.8. Cultures were induced with 1 mM isopropyl-$1$-thio-$\beta$-$D$-galactopyranoside and incubated at 18°C for 18 h. Cell pellets were frozen at $-80^\circ$C overnight and resuspended in 50 mM Tris–HCl buffer, pH 8.0 containing 0.5 M NaCl, 50 mM glycine, 5% (v/v) glycerol, 20 mM imidazole and protease inhibitors (complete EDTA-free tablets, one tablet per 50 ml, Roche). Cells were lysed by sonication. Histidine-tagged SFI3 was purified from cleared lysate using an ÄKTAxpress system (GE Healthcare) by IMAC on a 5 ml Ni$^{2+}$ His-Trap FF column followed by gel filtration on a Superdex 75 26/60 column pre-equilibrated in 20 mM HEPES, pH 7.5 containing 0.15 M NaCl (gel filtration buffer). Eluted monomeric protein was cleaved overnight at $4^\circ$C with histidine-tagged 3C protease (10 μg/mg protein) in gel filtration buffer supplemented with 5 mM β-mercaptoethanol. Cleaved SFI3 protein was separated from uncleaved protein by IMAC chromatography, concentrated to 5–7 ml and re-purified by gel filtration as above. Monomeric protein was concentrated to 16 mg ml$^{-1}$, frozen in 50 μl aliquots in liquid nitrogen and stored at $-80^\circ$C. SFI3 protein concentration was estimated by measuring absorbance at 280 nm and predicted extinction coefficients for the fusion and cleaved proteins of 19 940 M$^{-1}$ cm$^{-1}$.

**Crystallisation, data collection and structure solution**

For crystallisation, SFI3 was concentrated to 20 mg ml$^{-1}$. Crystallisation experiments used 96-well plates, commercially available reagents, and were set up with an Oryx Nano robot (Douglas Instruments). Optimizations were carried out in 48-well plates, also using an Oryx Nano robot. For X-ray data collection, crystals were grown in 0.1 M bis-tris-propane pH 7.5, 0.2 M sodium acetate and 20% (w/v) polyethylene glycol (PEG) 3350, transferred to a cryoprotectant solution containing 0.1M bis-tris-propane pH 7.5, 0.2 M sodium acetate and 35% (w/v) PEG 3350, mounted in a lithio-loop and flash cooled in liquid nitrogen. To enable collection of anomalous X-ray data for structure solution, crystals were soaked for c. 60 s in a well solution supplemented with 0.1 M potassium iodide and cryoprotected as above before flash freezing in liquid nitrogen.

Native and single wavelength anomalous diffraction (SAD) X-ray data sets were collected at the Diamond Light Source, United Kingdom, beamline I02 (under beamtime allocation MX1219). The data were processed using xia2 (Winter, 2010), and scaled with SCALA from the CCP4 suite (Winn et al., 2011). The structure was solved using SAD data collected from a crystal soaked in potassium iodide solution. Cocrystals were identified with Phenix, which was also used to calculate phases and generate initial electron density maps. The experimental phases were then used by ARP/wARP, as implemented in CCP4, to autobuild the first model. This model was subsequently used to phase a second crystal form by molecular replacement. The final model (in the latter crystal form) was produced through iterative rounds of refinement using REFMAC5 (Murshudov et al., 1997) and manual rebuilding with COOT (Emsley et al., 2010). Structure validation used the tools provided in COOT and MOLPROBITY (Chen et al., 2010a,b).

The co-ordinates and structure factors for SFI3 have been deposited at the Protein Data Bank (PDB) with accession number 6GU1.
Agrobacterium-mediated transient expression

Agrobacterium-mediated transient expression by agroinfiltration was performed as described previously (Gilroy et al., 2011; Yang et al., 2016; Boevink et al., 2016b). Agrobacterium strains were infiltrated into leaves of N. benthamiana wild-type or VIGS plants. The concentrations measured by absorbance at 600 nm of A. tumefaciens used were: 0.1–0.01 for confocal imaging, 0.5 for western blot analyses and HR assays and 0.1 for agroinfiltration and infection assays. For co-expression of multiple constructs A. tumefaciens suspensions of strains carrying the different constructs were thoroughly mixed before infiltration.

Confocal microscopy

Agrobacterium tumefaciens containing target protein fusions were infiltrated into leaves of 4-wk-old N. benthamiana plants. Nicotiana benthamiana leaf cells expressing fluorescent protein fusions were imaged no later than 2 d after agroinfiltration using a Nikon A1R confocal microscopes confocal microscope. GFP was excited with 488 nm from an argon laser and its emissions were detected between 500 and 530 nm. Images were collected from leaf cells expressing low levels of the protein fusions to minimise artefacts of ectopic protein expression. MG132 (40 μM) was infiltrated into leaves 6 h before imaging under the confocal microscope.

Yeast-two-hybrid

A Y2H screen with pDEST32-PiSFI3 was performed as described in McLellan et al. (2013) using the Invitrogen ProQuest system. The full-length coding sequence of the candidate interacting prey sequence (U-box kinase) was cloned and re-tested with pDEST32-PiSFI3 and use pDEST32-Pi04089 as a control to rule out the possibility that the observed reporter gene activation had resulted from interactions between the prey and the DNA-binding domain of the bait construct or DNA-binding activity of the prey itself. For each transformation, 1 μg plasmid DNA and 100 μg denatured sheared salmon sperm DNA were mixed together with 100 μl of the yeast suspension; 700 μl of 1× lithium acetate/40% PEG 3350/1× Tris–EDTA (TE) buffer were added, mixed well and incubated at 30°C for 30 min; 88 μl dimethyl sulphoxide (DMSO) were added, mixed well, heat shocked at 42°C for 7 min, centrifuged for 10 s and the supernatant was removed. The pellet was resuspended in 1 ml 1× TE and repelleted; the pellet was resuspended in 50-100 μl TE and plated on a selective media. Transformants were identified by testing reporter gene activation in the X-gal assay.

Co-immunoprecipitation analyses

The fused protein constructs were transiently overexpressed in N. benthamiana by Agrobacterium-mediated expression. Leaf samples were collected 48 h after infiltration. Proteins were extracted by using GTEN buffer (25 mM Tris–HCl pH 7.5, 1 mM EDTA, 150 mM NaCl, 10% glycerol) with 10 mM dithiothreitol (DTT), protease inhibitor cocktail (PIC), 1 mM pregnant mare’s serum gonadotropin (PMSF) and 0.2% Nonidet P40. Samples were ground in liquid nitrogen with 500 μl extraction buffer on ice, gently vortexed and placed on ice and centrifuged at 13 000 g for 10 min at 4°C; the supernatant was transferred to a precooled tube (re-spin if leaf tissues were present in the supernatant, until clear); To 50 μl sample (input) 50 μl 2× sample buffer were added and boiled at 95°C for 10 min and stored at −20°C for western blot analysis; 450 μl of extract was kept for the co-immunoprecipitation (Co-IP); 20–25 μl beads were used per immunoprecipitation (IP). The required amount was taken and washed three times with ice-cold dilution/wash buffer; washed beads were added to cell lysates and shaken for 1–2 h at 4°C or room temperature (RT). Beads were washed twice with 500 μl ice cold wash buffer, resuspended in 50 μl 2× sodium dodecyl sulphate (SDS) buffer and boiled at 95°C for 10 min. The samples were stored at −20°C for western analysis. mRFP-tagged or GFP-tagged fusions were immunoprecipitated using mRFP-Trap®-M magnetic beads or GFP-Trap®-M magnetic beads (Chromotek Gmbh), respectively. The resulting samples were separated by SDS-PAGE and western blotting. Immunoprecipitated RFP fusions and co-immunoprecipitated c-MYC fusions were detected using appropriate antisera.

Flg22 treatment and gene expression assays

Here, 6-wk-old potato plants or 3-wk-old VIGS plants were used for the treatments. Three leaves from the third to the fifth compound leaf from the top of each potato line or three leaves from VIGS-plants were infiltrated with flg22 (40 mM). Three leaves per time point each were harvested from separate plants and snap frozen in liquid nitrogen. The RNA was extracted and cDNA was made and diluted to 1:20 for RT-PCR using the marker genes. Primer pairs were designed outside the region of cDNA targeted for silencing following the manufacturer’s guidelines. Primer sequences are given in Table S1. Relative expression of the target genes was calculated using the 2ΔΔCt method with StTubulin as the reference for potato, Nhef1α for N. benthamiana. Actin was used as the reference of potato for semi-quantified RT-PCR.

The P. infestans biomass was quantified by qPCR using Ef-1α (used to quantify S. tuberosum DNA) and PiO8 primers (designed based on highly repetitive sequences from the P. infestans genome to quantify P. infestans DNA) as described by Llorente et al. (2010). Three leaves for each time point (every 24 h after inoculation with P. infestans) were harvested and snap frozen in liquid nitrogen. The DNA was extracted and quantified by qRT-PCR.

Results

Nuclear localization of GFP–PiSFI3 is important for P. infestans virulence

Transient expression of GFP–PiSFI3 in N. benthamiana showed that it was enriched in the nucleus/nucleolus and enhanced
colonization by *P. infestans* (Zheng et al., 2014). To investigate whether the nucleus is an important site of PiSFI3 activity within host cells, we misdirected its localization. In addition to GFP–PiSFI3, two fusion constructs were generated, either with a nuclear export signal (NES; NESGFP–PiSFI3) or with a nuclear localization signal (NLS; NLSGFP–PiSFI3), as described for other RXLR effectors (Boevink et al., 2016b; Yang et al., 2016). As shown previously (Zheng et al., 2014, 2018), GFP–PiSFI3 accumulated strongly in the nucleus, forming a ring around the nucleolus, with additional cytoplasmic background (Fig. 1a). By contrast, NESGFP–PiSFI3 fluorescence was observed primarily in the cytoplasm, whereas NLSGFP–PiSFI3 fluorescence was focussed in the nucleus and nucleolus (Fig. 1a). GFP–PiSFI3, NLSGFP–PiSFI3 and NESGFP–PiSFI3 fusions were expressed transiently in *N. benthamiana* and the leaves were challenged with *P. infestans* 1 d after agroinfiltration. Lesion size was recorded after 6 d. As described in Zheng et al. (2014, 2018), GFP–PiSFI3 enhanced colonization by *P. infestans*. Expression of NLSGFP–PiSFI3 was found to also significantly promote larger *P. infestans* lesions compared with free GFP control (analysis of variance [ANOVA], *P* < 0.001). Lesion sizes following the expression of NESGFP–PiSFI3 were reduced significantly compared with GFP–PiSFI3 and NLSGFP–PiSFI3, but showed no significant difference compared to the empty GFP control (Fig. 1b, Supporting Information Figure S1). All constructs produced intact fusion proteins when expressed *in planta* (Fig. 1c). We therefore conclude that exclusion of PiSFI3 from the nucleus attenuates its contribution to virulence.

**PiSFI3 attenuates early flg22-triggered transcriptional responses in potato**

To determine whether PiSFI3 enhanced *P. infestans* colonization in potato, cultivar E-potato-3 (E3) was transformed with a construct expressing 35S:PiSFI3. Six independent transgenic lines were selected for the preliminary experiments. Following inoculation of zoospores, pathogen colonisation was considerably enhanced (*P* < 0.001, one-way ANOVA) on the transgenic lines compared with the E3 control, as measured by lesion diameter at 5 d post-inoculation (dpi; Fig. S2). The transgenic lines expressing SFI3 showed no developmental phenotype, and were similar to the E3 control. Two transgenic lines, PiSFI3-25 and
PiSFI3-26, were selected for further studies (Fig. 2a). These two lines showed significantly enhanced *P. infestans* colonization ($P < 0.001$, one-way ANOVA) compared with the E3 control, as measured by lesion diameter and sporangia number (Figs 2b–d, S2c). Moreover, *P. infestans* biomass, determined by qPCR measurement of pathogen DNA, was observed to increase more rapidly on the transgenic PiSFI3-25 and PiSFI3-26 lines between 2 and 3 dpi, which is the biotrophic phase of infection before symptom development (Fig. 2e). These results confirmed previous observations that transient expression of SFI3 on the model *P. infestans* host *Nicotiana benthamiana* also enhanced pathogen colonisation (Zheng et al., 2014, 2018).

PiSFI3 was previously shown to act downstream of MAP kinase activation in suppressing pFRK:luc reporter gene induction following flg22 treatment (Zheng et al., 2014). To determine whether PiSFI3 suppresses early flg22-mediated transcriptional responses on potato, two flg22 marker genes, *NbWRKY7* and *NbACRE31*, used previously in *N. benthamiana* (McLellan et al., 2013), were selected and their expression in potato E3 assessed after flg22 treatment. Transcript accumulation of each gene was shown to peak as early as 30 min posttreatment with flg22 (Fig. S3), and this time point was therefore selected to study the transgenic lines. Following treatment of the transgenic lines and E3 control with flg22 peptide, transcript accumulation of both *StWRKY7* and *StACRE31* (Figs 2f, g, S3c, d) was attenuated in the transgenic PiSFI3 lines compared to the E3 control, demonstrating that the effector also attenuates early flg22-mediated transcriptional responses in potato leaves.

**Fig. 2** PiSFI3 attenuates early flg22-triggered transcriptional responses in potato. (a) Gel images of a semi-quantitative PCR to compare PiSFI3 and control Actin transcripts amplified from cDNA made from control and transgenic potato cultivar E3 (cv). (b) Representative trypan blue-stained leaf images of leaves of transgenic potatoes cv E3 expressing PiSFI3 and E3 controls infected with a mixture of two *Phytophthora infestans* isolates (HB09-14-2 and HB-0916-2) and photographed at 5 d post inoculation (dpi). (c) Boxplots shows the mean (red line) and median (black line) *P. infestans* lesion diameter of cv E3 and transgenic PiSFI3-expressing potato plants measured at 5 dpi. Boxplots represent the combined data from three biological reps (46 leaves per line). Letters on the boxplots denote statistically significant differences (ANOVA, $P < 0.001$). The black dots above and below each box represent the 95th and 5th percentage of outliers, respectively. (d) Overexpressing PiSFI3 allow a significant increase in *P. infestans* sporulation compared to the E3 control. Boxplots represent the combined data from three biological reps ($n = 120$). Letters on the boxplots denote statistically significant differences (ANOVA, $P < 0.001$). The black line represents the median line and the red line is the mean. The black dot above and below each box represent the 95th and 5th percentage of outliers. (e) Graph shows *P. infestans* biomass, determined by qPCR measurement of pathogen DNA using Ef-1a (used to quantify *Solanum tuberosum* DNA) and PiO8 primers (designed based on highly repetitive sequences from the *P. infestans* genome to quantify *P. infestans*). Three leaves for each time point (every 24 h after inoculation with *P. infestans*) were harvested and snap frozen in liquid nitrogen. The DNA was extract and quantified by qRT-PCR. Error bars show ± SE. (f, g) Relative expression of flg22 marker genes *StWRKY7* and *StACRE31* 30 min after treatment with flg22 in cv E3, PiSFI3-25 and PiSFI3-26 lines. Error bars show ± SE.
PiSFI3 does not suppress programmed cell death (PCD) triggered by INF1 or Cf4/Avr4

In the previous study by Zheng et al. (2014, 2018), the three effectors that suppress flg22-mediated posttranslational MAP kinase activation (PiSFI5, PiSFI6, PiSFI7) were tested to see if they suppress two independent signal transduction pathways, triggered by expression of the elicitin infestin-1 (INF1), or by co-expression of tomato Cf4 with Cladosporium fulvum Avr4, which led to PCD. Whereas AVR3a was able to strongly suppress both INF1- and Cf4/Avr4-mediated PCD, only weak suppression was observed with PiSFI7, and only of INF1-mediated PCD (Zheng et al., 2014, 2018). We therefore extended the analysis to PiSFI3. Again, whereas both INF1- and Cf4/Avr4-mediated PCD were strongly suppressed by the positive control, AVR3a, no such suppression was observed with PiSFI3 (Fig. S4). This result indicates that PTI suppression by PiSFI3 is restricted, in that it functions to attenuate flg22-mediated early transcriptional events but does not attenuate INF1-mediated PCD or, indeed, another pathway leading to PCD, triggered by Cf4 activation.

PiSFI3 interacts with a potato protein containing U-box and kinase domains

To investigate what PiSFI3 targets in the host to suppress flg22-mediated responses, a yeast-2-hybrid (Y2H) library of cDNA made from potato infected with P. infestans (Bos et al., 2010) was screened with a GAL4 DNA binding domain–PiSFI3 fusion (‘bait’) construct to a depth of 2.6 × 10^6 yeast co-transformants. Twelve independent yeast colonies recovered from selection plates that contained GAL4 activation domain (‘prey’) fusions yielded sequences encoding a potato protein containing both U-box and kinase domains. The gene was therefore called Solanum tuberosum U-box and kinase domain (StUBK), and is a reciprocal best BLAST hit (candidate orthologue) of the gene AtPUB33 in Arabidopsis thaliana (Fig. S5), the function of which is unknown. The plant U-box (PUB) domain is an ubiquitin E3 ligase domain and is uniquely combined with a kinase domain in AtPUB33. To confirm the Y2H interaction, a full-length StUBK sequence was cloned into the prey construct and tested pairwise with bait constructs containing PiSFI3, along with a control RXLR effector (Pi04089) which shares a similar nuclear and nucleolar localisation in plant cells, and which targets the RNA binding protein StKRBP1 (Wang et al., 2015) and an empty bait vector (EV).

Whereas all transformants grew on the control plates (+HIS) only yeast cells containing both PiSFI3 and StUBK were able to grow on the selection (−HIS) plates and activate the β-galactosidase (β-Gal) reporter (Fig. 3a).

To confirm that PiSFI3 and StUBK also interact in planta, co-immunoprecipitation (Co-IP) experiments were performed using cMYC–PiSFI3 and N-terminal mRFP-tagged StUBK (mRFP–StUBK), with cMYC–GUS and mRFP–GUS used as controls. The mRFP and cMYC fusion constructs were transiently co-
expressed in *N. benthamiana*. While all constructs were detected in the relevant input fractions, only cMYC–PiSFI3 and not the cMYC–GUS control was co-immunoprecipitated on mRFP–TRAP_M beads in the presence of mRFP–StUBK, and mRFP–GUS was unable to co-immunoprecipitate cMYC–PiSFI3 (Fig. 3b). As an additional control, we demonstrated that, by contrast to PiSFI3, the nuclear effector control Pi04089 failed to interact with mRFP–StUBK in planta (Fig. S6a).

To investigate the subcellular localization of StUBK, GFP was fused to its N terminus to form GFP–StUBK and viewed following *Agrobacterium*-mediated expression in *N. benthamiana* using confocal microscopy. GFP–StUBK protein localized in the nucleus with a ring around the nucleolus, and showed additional cytoplasmic fluorescence background (Fig. 3c), which is a similar localization to PiSFI3 (Fig. 3c). GFP–StUBK was stable as a fusion protein in planta (Fig. 3d).

**Silencing of UBK enhances *P. infestans* leaf colonisation**

To investigate a possible contribution of UBK to immunity, VIGS was used to knock-down *NbUBK* transcript levels in *N. benthamiana*. *Nicotiana benthamiana* is an alternative host for *P. infestans* and serves as a model for studying late blight, as it allows transient *Agrobacterium*-mediated gene expression, VIGS and noninvasive confocal microscopy to aid functional studies of molecular interactions between host and pathogen (e.g. Bos et al., 2010; McLellan et al., 2013; King et al., 2014; Zheng et al., 2014, 2018; Boevink et al., 2016b; Murphy et al., 2018).

Two independent TRV constructs (UBK-V1 and UBK-V2) were generated to specifically silence *NbUBK* by cloning two portions of the gene upstream of the conserved U-box and kinase domains to avoid off-target silencing (Fig. S7a). qRT-PCR was used to test silencing levels in each of three independent biological replicates and showed that transcript levels of *NbUBK* were reduced by 60–90% (Fig. S7b). *Nicotiana benthamiana* plants expressing UBK-V1 or UBK-V2 constructs consistently showed increased *P. infestans* colonisation compared to control TRV-GFP plants, measured as the percentage of inoculation sites that developed lesions (*P* < 0.001, one-way ANOVA) and by counting the numbers of sporangia developing on leaves at 8 dpi (*P* < 0.002, one-way ANOVA; Fig. 4a–c). This indicates that silencing *NbUBK* leads to increased susceptibility.

To investigate whether silencing *StUBK* has a similar effect on pathogenesis in potato, an antisense construct was generated, using the equivalent portion of the *StUBK* gene that was cloned from *NbUBK* in order to generate TRV construct UBK-V2 (Fig. S7c). This construct was transformed into potato cv E3. Six RNAi lines generated in the preliminary experiments revealed an 80% reduction in *StUBK* transcript accumulation (Fig. S7d). Each of the RNAi lines showed similar, significant enhancement of *P. infestans* colonisation (*P* < 0.001, one-way ANOVA; Fig. S8a), which was apparent using trypan blue (Fig. S8b). Three of these RNAi lines were selected (i2-24, i2-32, i2-39) for further analysis as they showed 80–90% reduction in *StUBK* transcript accumulation (Fig. S7d). *Phytophthora infestans* colonisation, measured as *P. infestans* biomass, determined by qPCR measurement of pathogen DNA, was significantly (*P* < 0.001, one-way ANOVA) more extensive between 2 and 3 dpi compared with the untransformed cultivar E3 (measured for line i2-39 in Fig. 4d). Moreover, all three lines showed significantly enhanced *P. infestans* colonisation, as measured by lesion diameter and number of sporangia (Figs 4e,f, S8b), by stained *P. infestans* mycelia (Bos et al., 2010).

**Overexpression of STUBK reduces *P. infestans* leaf colonisation**

We investigated the affect of overexpression of *StUBK* on late blight disease. Critically, transient expression of GFP–StUBK led to significantly (*P* < 0.05, one-way ANOVA) reduced *P. infestans* colonisation compared with the GFP control (Fig. 4g). Taken together, the VIGS in *N. benthamiana*, RNAi in potato and overexpression demonstrate that UBK contributes to restricting *P. infestans* infection.

To examine whether overexpression of *StUBK* has an effect on the virulence activity of PiSFI3, GFP–PiSFI3 was transient expressed in *N. benthamiana* alone or co-expressed with GFP–StUBK and challenged with *P. infestans* spores at 1 dpi, using GFP–EV and GFP–Pi04089 as controls. At 6 dpi, GFP–PiSFI3 or GFP–Pi04089 expressed alone enhanced *P. infestans* colonisation to a significant level (*P* < 0.001, one-way ANOVA) compared with the GFP–EV control (Fig. 4h), as anticipated. However, the ability of GFP–PiSFI3 to enhance *P. infestans* colonisation was significantly reduced (ANOVA, *P* < 0.5) by co-expression with GFP–StUBK. By contrast, enhanced colonisation of *P. infestans* by GFP–Pi04089 was unaltered when co-expressed with GFP–StUBK (Fig. 4h). STUBK overexpression was therefore able to specifically reduce PiSFI3 virulence activity.

**VIGS of UBK does not compromise INF1 or CF4/Avr4 cell deaths**

PiSFI3 attenuates flg22-mediated early transcriptional events, but does not suppress PCD triggered by INF1 or CF4/Avr4. We therefore proceeded to investigate whether PCD triggered in *N. benthamiana* by measurement of INF1, or if CF4/Avr4 requires UBK. Silencing of UBK did not attenuate INF1 or CF4/Avr4 cell death, indicating that UBK is not involved in the cell death responses to INF1 and Avr4 (Fig. S9).

**Silencing of UBK attenuates early flg22 induced marker genes**

PiSFI3 suppressed pFRK:Luc reporter induction following flg22 treatment (Zheng et al., 2014, 2018) and suppressed flg22-mediated early gene expression (*NbWRKY7* and *NbACRE3i*) in potato (Fig. 2f,g). We investigated whether the expression of *StUBK* also responded to the flg22 treatment. Following flg22 infiltration into leaves (performed as described previously by He et al., 2015 and McLellan et al., 2013), increased *StUBK* transcript accumulation was noted as early as 3 h postinoculation,
Fig. 4 Silencing of UBK enhances, and overexpression reduces, Phytophthora infestans leaf colonisation. (a) Representative leaf images show the extent of *P. infestans* leaf colonisation on plants expressing each two independent virus-induced gene silencing (VIGS) constructs (UBK- V1 and UBK- V2) in *Nicotiana benthamiana*. (b) Silencing of UBK using two independent VIGS constructs (UBK-V1 and UBK- V2) in *N. benthamiana* significantly increases *P. infestans* lesion number compared with the GFP control. Boxplots represent the combined data from five biological reps. Letters on the boxplots denote statistically significant differences (ANOVA, *P* < 0.001). (c) Silencing of UBK using two independent virus-induced gene silencing (VIGS) constructs allow a significant increase in *P. infestans* sporulation compared with the GFP control. Boxplots represent the combined data from four biological reps (*n* = 32). Letters on the boxplots denote statistically significant differences (ANOVA, *P* < 0.05). (d) Graph shows that silencing STUBK by RNAi in potato significantly increases *P. infestans* biomass compared to the GFP control. *Phytophthora infestans* biomass determined by qPCR measurement of pathogen DNA using *Ef-1*α (used to quantify *Solanum tuberosum* DNA) and PiO8 primers (designed based on highly repetitive sequences from the *P. infestans* genome to quantify *P. infestans*). Three leaves for each time point (every 24 h after inoculation with *P. infestans*) were harvested and snap frozen in liquid nitrogen. The DNA was extract and quantified by qRT-PCR. (e) Boxplots show mean *P. infestans* lesion diameter of potato cultivar E3 and transgenic STUBK RNAi plants measured at 5 d post inoculation (dpi). Boxplots represent the combined data from three biological reps (*n* = 34 leaves per line). Letters on the boxplots denote statistically significant differences (ANOVA, *P* < 0.001). (f) Transgenic potato RNAi lines silencing STUBK allow a significant increase in *P. infestans* sporulation compared to the E3 control. Boxplots represent the combined data from three biological reps (*n* = 96). Letters on the boxplots denote statistically significant differences (ANOVA, *P* < 0.001). (g) GFP–StUBK following Agrobacterium-mediated expression significantly (*P* < 0.05) reduces the ability of *P. infestans* to colonise *N. benthamiana* compared with the GFP–EV control. Lesion size was recorded after 6 d. Boxplots represent the combined data from three biological reps (*n* = 78). (h) GFP–PiSFI3 and GFP–Pi04089 following Agrobacterium-mediated expression significantly (*P* < 0.001) enhance *P. infestans* colonization compared to GFP–EV control. GFP–StUBK significantly (*P* < 0.05) reduces the ability of GFP–PiSFI3 but not GFP–Pi04089 to colonise *N. benthamiana* compared to GFP–EV control or GFP–PiSFI3. Lesion size was recorded after 6 d. Boxplots represent the combined data from three biological reps (*n* = 60). For each boxplot the black line represents the median and the red line the mean. The black dot above and below each box represent the 95th and 5th percentage of outliers, respectively.
peaking at 6 h (Fig. S10a). This prompted us to investigate whether UBK contributes to the flg22-mediated PTI response, using the early-induced genes NbWRKY7 and NbACRE31 as transcriptional markers. Following flg22 treatment of NbUBK VIGS plants, NbWRKY7 and NbACRE31 transcript accumulation was attenuated compared with the TRV-GFP control (Fig. 5a,b). Similarly, a reduced accumulation of StWRKY7 and StACRE31 transcripts was observed in the three RNAi lines of StUBK (i2-24, i2-32, i2-39) compared with the untransformed potato cultivar E3 control (Figs 5c,d, S10b,c). This indicates that UBK contributes to the flg22-triggered PTI transcriptional response.

**PiSFI3 is a WY-domain RXLR effector with a trans-homodimer fold**

To investigate how PiSFI3 may function at the molecular level, we determined its crystal structure to 1.7 Å resolution (Table S2). The PiSFI3 structure comprises residues Ala63–Lys117 of the full-length sequence and adopts an antiparallel extended α-helical fold (Fig. 6a). The structure confirms that SFI3 adopts the ‘WY-domain’ fold, as previously predicted (Boutemy et al., 2011). However, unlike previous structures of RXLR effectors, PiSFI3 forms a novel ‘trans’ WY-domain configuration in which the ‘W’ (Trp) and ‘Y’ (Tyr) residues are derived from separate monomers (Fig. 6a). Therefore, the SFI3 structure further highlights the adaptability of the WY-domain to form diverse conformations (Winn et al., 2011; Maqbool et al., 2016, Fig. S11a). To test whether PiSFI3 can oligomerise in planta we used Co-IP with cMYC- and mRFP-tagged versions of the protein co-expressed in N. benthamiana (Fig. 6b). Following immunoprecipitation with anti-mRFP antibodies we detected cMYC–PiSFI3 protein in samples where mRFP–PiSFI3 was co-expressed, but not the mRFP–GUS control, confirming that PiSFI3 can oligomerise in plant cells (Fig. 6b).

We then examined the PiSFI3 structure to identify amino acid residues that, on mutation, might perturb protein structure and inform functional studies. For this we chose to generate the double mutants PiSFI3<sup>L86D/L100D</sup> and PiSFI3<sup>L86D/L103D</sup>. These pairs of residues faced each other across the PiSFI3 α-helices and Leu/Asp mutations were predicted to result in conformational changes in the protein, potentially disrupting oligomerisation (Fig. 6a). Indeed, whereas mRFP–PiSFI3 co-immunoprecipitated cMYC–PiSFI3, co-immunoprecipitation of cMYC–PiSFI3<sup>L86D/L100D</sup> and cMYC–PiSFI3<sup>L86D/L103D</sup> by, respectively, mRFP–PiSFI3<sup>L86D/L100D</sup> and mRFP–PiSFI3<sup>L86D/L103D</sup> was attenuated (Figs 6b, S11b).

**Structure-guided mutations of SFI3 are attenuated in virulence function and lose nucleolar localisation**

As shown above, nuclear/nucleolar localization of GFP–PiSFI3 is important for P. infestans virulence. To determine the subcellular localization of GFP–PiSFI3<sup>L86D/L100D</sup> and GFP–PiSFI3<sup>L86D/L103D</sup>, we transiently expressed these mutants in N. benthamiana alongside GFP–PiSFI3 as a positive control. As previously observed, GFP–PiSFI3 accumulated strongly in the nucleus, forming a ring around the nucleolus, with additional cytoplasmic background (Fig. 7a). By contrast, GFP–PiSFI3<sup>L86D/L100D</sup> and GFP–PiSFI3<sup>L86D/L103D</sup> fluorescence was primarily in the nucleoplasm, with cytoplasmic background, and was reduced in the nucleolus, leading to a

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**Fig. 5** Silencing of UBK attenuates early flg22 induced marker genes. (a, b) Relative expression of flg22 marker genes NbWRKY7 and NbACRE31 3 h after treatment with flg22 in GFP, and UBK VIGS plants (UBK-V1 and UBK-V2). (c, d) Relative expression of flg22 marker genes StWRKY7 and StACRE31 30 min after treatment with flg22 in E3, StUBK RNAi lines (i2-24, i2-32, i2-39). Error bars represent ± SE. RNA was extracted to synthesis cDNA and qRT-PCR reactions were performed using Power SYBR Green for gene expression assays. Primers used are available in Supporting Information Table S1. Relative expression of the target genes was calculated using the 2<sup>-ΔΔCT</sup> method with the STUBI housekeeping gene as the reference for potato as described previously (Livak & Schmittgen, 2001; McLellan et al., 2013).
significantly reduced ratio of nucleolar-to-nucleoplasmic fluorescence (Figs 7a,b, S12). GFP–PiSFI3, GFP–PiSFI3L86D/L100D and GFP–PiSFI3L86D/L103D fusions were stable in planta (Fig. S12). As expression of PiSFI3 in planta enhances infection (Fig. 1), we tested whether the mutations altered effector virulence function. GFP–PiSFI3, GFP–PiSFI3L86D/L100D and GFP–PiSFI3L86D/L103D were transiently expressed in one-half of an N. benthamiana leaf with empty GFP expressed as a control in the other. The two halves of the leaves were drop-inoculated with P. infestans zoospores and lesion size was measured at 6 d postinoculation. Unlike the wild-type GFP–PiSFI3, both GFP–PiSFI3L86D/L100D and GFP–PiSFI3L86D/L103D failed to enhance P. infestans colonization, and each showed no significant difference compared with a GFP–EV control (ANOVA, \( P < 0.001 \); Figs 7c, S12c).

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**Fig. 6** SFI3 forms a homodimer with an unusual ‘trans-WY’ configuration and self-associates in planta. (a) Overall schematic view of the crystal structure of SFI3. SFI3 comprises two extended helical folds (monomers coloured blue and orange) that form a dimer in a ‘trans-WY’ configuration. The positions of the N- and C-termini, Trp72 and Tyr107, and of the Leu residues (Leu86, Leu100 and Leu103) that were mutated to disrupt dimer formation, are labelled in each monomer and shown with side chains in stick representation. Structure figures were prepared with CCP4MG (McNicholas et al., 2011). (b) SFI3 can oligomerise in planta by Co-IP. cMYC- and mRFP-tagged versions of the proteins co-expressed in Nicotiana benthamiana. Following immunoprecipitation with anti-mRFP antibodies, cMYC-tagged PiSFI3 protein in samples where mRFP–PiSFI3 has been expressed, but not the mRFP–GUS control, confirming PiSFI3 can oligomerise in plant cells. Co-immunoprecipitation of cMYC–PiSFI3L86D/L100D and cMYC–PiSFI3L86D/L103D by, respectively, mRFP–PiSFI3L86D/L100D and mRFP–PiSFI3L86D/L103D was attenuated. Expression of constructs in the leaves is indicated by +. Protein size markers are indicated in kDa, and protein loading is indicated by Ponceau stain (PS).
PiSFI3 mutations prevent interaction with StUBK

As PiSFI3 mutations showed reduced virulence activity, this prompted us to investigate whether the mutations had an effect on the interaction with StUBK. StUBK was cloned into the Y2H prey construct and tested pairwise with bait constructs containing PiSFI3, PiSFI3L86D/L100D and PiSFI3L86D/L103D along with a control RXLR effector (Pi04089). Whereas all transformants grew on the control plates (+HIS), only yeast containing both PiSFI3 and StUBK was able to grow on the selection (-C0HIS) plates and activate the β-galactosidase (β-GAL) reporter (Fig. 8a). The result indicates that PiSFI3L86D/L100D and PiSFI3L86D/L103D failed to interact with StUBK in a Y2H assay.

To further investigate the interaction between StUBK and the two mutants in planta, RFP-tagged StUBK was used in co-immunoprecipitation experiments. cMYC–PiSFI3, cMYC–PiSFI3L86D/L100D and cMYC–PiSFI3L86D/L103D were co-expressed in N. benthamiana with N-terminal RFP-tagged StUBK, respectively, with cMYC–GUS used as a noninteracting control. All constructs were detected in the relevant input fractions. Following immunoprecipitation using RFP–TRAP_M beads, only the cMYC–SFI3 construct, and not the cMYC–PiSFI3L86D/L100D or cMYC–PiSFI3L86D/L103D, was detected in the presence of RFP-UBK (Fig. 8b), confirming that PiSFI3L86D/L100D and PiSFI3L86D/L103D fail to interact with StUBK in planta.

Discussion

The RXLR effector PiSFI3 acts downstream of MAP kinase activation to suppress flg22-triggered transcriptional defence responses (Zheng et al., 2014, 2018). Here we show that PiSFI3 forms a trans-homodimer that targets a plant protein that uniquely contains both U-box and kinase domains (StUBK). StUBK is a positive regulator specifically of early transcriptional changes that can be triggered by the bacterial MAMP flg22.

Transient expression of GFP–PiSFI3 in N. benthamiana enhanced colonization by P. infestans (Zheng et al., 2014, 2018). Although future work to silence expression of PiSFI3 in P. infestans may reveal whether this effector is essential for infection, we
demonstrate that its stable transgenic expression in potato resulted in enhanced *P. infestans* leaf colonization, supporting its role as an effector (Fig. 2). Moreover, transgenic PiSFI3-expressing potato lines attenuate early flg22-mediated transcriptional gene expression in agreement with the previous assay reported in tomato protoplasts (Zheng *et al.*, 2014, 2018). As *P. infestans* does not possess the MAMP flagellin, the ability of PiSFI3 to attenuate flg22-triggered early defences indicates that the receptor FLS2 activates a generic signal transduction pathway that is likely to be also activated by an unknown receptor that detects an oomycete MAMP (Zheng *et al.*, 2014, 2018). It will be interesting to identify both the host receptor(s) and *P. infestans* elicitor(s) responsible for activating this defence pathway in future work.

Suppression of flg22-triggered transcriptional responses by PiSFI3 is mediated by its interaction with StUBK. We show that silencing *UBK* in both potato and *N. benthamiana* attenuates flg22-triggered transcriptional responses (Fig. 5). This silencing leads to increased *P. infestans* susceptibility, whereas overexpression of StUBK increases resistance to *P. infestans* (Fig. 4). Critically, co-expression of StUBK with either PiSFI3 or the control effector Pi04089 specifically attenuated the virulence activity of the former (Fig. 4), providing evidence that UBK is the likely operative target of SFI3. Further genetic evidence indicating this to be the case was provided by mutation of PiSFI3 that linked its ability to interact with StUBK with its ability to enhance *P. infestans* colonisation (Figs 7, 8). Interestingly, PiSFI3 overexpression or UBK silencing had no impact upon cell death triggered by perception of the *P. infestans* MAMP INF1 (Figs S4, S9), suggesting that effector and target are specific to a particular defence pathway. Future work will investigate whether INF1 activates early transcriptional responses in common with flg22 perception that led to defences other than cell death, and which are suppressed by PiSFI3.

Fig. 8 Structure-guided mutation of SFI3 prevents its interaction with UBK. (a) Yeast co-expressing StUBK with PiSFI3 grew on –histidine (–HIS) medium and yielded β-galactosidase (β-Gal) activity, while those co-expressed with the control Pi04089, PiSFI3L86D/L100D, or PiSFI3L86D/L103D did not. The +HIS control shows all yeast were able to grow in the presence of histidine. (b) These interactions were further confirmed by co-immunoprecipitation (Co-IP) assay using protein extracted from agroinfiltrated *Nicotiana benthamiana* leaves. cMYC–PiSFI3 associated with mRFP–StUBK, whereas cMYC–GUS, cMYC–PiSFI3L86D/L100D, or cMYC–PiSFI3L86D/L103D did not. Expression of constructs in the leaves is indicated by +. Protein size markers are indicated in kDa, and protein loading is indicated by Ponceau stain (PS). The arrowheads show the RFP-tagged fusion proteins.
PiSFI3 localizes to the host nucleus, forming a ring around the nucleolus, with additional cytoplasmic background. The virulence activity of the effector was considerably reduced when it was directed away from the nucleus with a nuclear export signal (NES), suggesting that the nucleus is an important site of PiSFI3 action. In agreement, when PiSFI3 was focused in the nucleus with an NLS, the beneficial activity of the effector was not diminished (Fig. 1). While background levels of the NLS-GFP–PiSFI3 fusion may yet remain in the cytoplasm, we argue that the lack of a statistically significant change to enhanced *P. infestans* colonization by the effector suggests that a cytoplasmic stage is not critical for PiSFI3 function. We therefore conclude that PiSFI3 activity is primarily in the nucleus. This is further supported by two observations. Firstly, StUBK was also present in the nucleolus, again forming a ring around this nuclear structure (Fig. 3). Moreover, mutations of PiSFI3 that disrupt its dimerization, prevent its interaction with StUBK, and attenuate its virulence function, also result in its failure to accumulate in the nucleolus (Fig. 7). Whilst these mutant forms still reside in the nucleoplasan, their failure to form a ring around the nucleolus or accumulate within it, strongly implicates this as the primary site of activity. It is interesting to note that both CMPG1 (Gilroy et al., 2011) and PUB17 (He et al., 2015) accumulate also in the nucleolus. Future work will reveal more about the function of StUBK in the nucleus by combining mutations of the U-box and kinase domains in the protein with studies to mis-direct it away from the nucleus or to exclusively localise it there using an NLS.

PiSFI3 adopts the ‘WY-domain’ fold, as previously predicted (Boutemy et al., 2011). However, unlike previous structures of RXLR effectors, SFI3 forms a novel ‘trans’ WY-domain configuration that can oligomerise *in vitro*. We demonstrate that PiSFI3 also oligomerises *in planta*. Mutation of Leu/Asp pairs of residues (L86D/L100D, L86D/L103D) that face each other across the PiSFI3 α-helices resulted in conformational changes that disrupted oligomerisation (Fig. 6). Interestingly, we found that the two mutants (PiSFI3<sub>L86D/L100D</sub> and PiSFI3<sub>L86D/L103D</sub>) that were stable *in planta* but no longer accumulated in the nucleolus, interacted with StUBK, or enhanced *P. infestans* colonisation. This demonstrates that the trans-homodimer is biologically relevant, and highlights the flexibility of the WY-domain fold.

Plant kinases and E3 ubiquitin ligases are well known for their functions in a variety of stress responses. Both phosphorylation and ubiquitination can either positively or negatively regulate immunity (Mithoe & Menke, 2018; Trujillo, 2018). StUBK possesses both a kinase domain and a U-box domain, indicating that it potentially regulates immunity via these PTMs. Both kinases and E3 ligases are targeted by pathogen effectors, either to prevent their activity, or potentially to utilise it. For example, PexRD2 suppresses the kinase signalling activity of its target MAP3K (King et al., 2014) to prevent phosphorylation of its substrates. More recently, *P. infestans* effector Pi17316 has been shown to target the MAP3K StVIK, utilising or promoting its activity as an S factor to suppress immunity (Murphy et al., 2018). Similarly, whereas the *P. infestans* effector AVR3a targets the E3 ligase CMPG1 to prevent its normal function (Bos et al., 2010), and effector Avr-Piz-t from *Magnaporthe oryzae* directly interacts with E3 ligases APIP6 and APIP10 to inhibit their positive regulation of PTI (Park et al., 2012, 2016), the effector Pi02860 from *P. infestans* utilizes host susceptibility factor E3 ligase NRL1 to degrade the immune regulator SWAP70 (Yang et al., 2016; He et al., 2018). Presumably, PiSFI3 prevents the normal activity of StUBK. Indeed, this is supported by observation that the effector attenuates the specific immune pathway that is activated by StUBK. Further work is needed to determine whether PiSFI3 inhibits either phosphorylation or ubiquitination activities, or interferes with substrate specificity of StUBK.

There is considerable cross-talk between ubiquitination and phosphorylation in the regulation of immunity. Phosphorylation often serves as a marker that triggers subsequent ubiquitination, in particular when ubiquitination leads to degradation. In other cases, ubiquitination provide a switching mechanism that can turn on/off the kinase activity of certain proteins. One recurrent theme for PUB E3 ligases is their interaction with kinase domains (Trujillo, 2018). For example, the kinase FLS2 is a target for ubiquitination and turnover by PUB13 (Lu et al., 2011). In addition, it was reported that U-box E3 ligase PUB13 regulates the abundance of chitin receptor LYK5 protein (Lysine Motif-Resceptor-Like Kinase 5) that is required for the activation of the CERK1 intracellular kinase domain and induction of plant innate immunity (Cao et al., 2014; Liao et al., 2017). Moreover, some positive regulators of immune pathways that act at the transcriptional level, such as the JA regulator MYC2, are both activated by phosphorylation and immediately inactivated by ubiquitination (Zhai et al., 2013). Turnover of phosphorylated NPR1 is required for full induction of target genes and establishment of systemic acquired resistance (SAR; Spoel et al., 2009). It is therefore perhaps unsurprising that StUBK, a positive regulator of immunity at the transcriptional level, contains both E3 ligase and kinase domains. Understanding how these two major PTMs interact to regulate signal transduction is an important topic in characterization of StUBK function in plant immunity. Future work will aim to uncover the details surrounding the interplay between phosphorylation and ubiquitination in StUBK and how this regulation is controlled upon *P. infestans* perception in the regulation of immune gene transcription. Of particular interest will be the identification of substrates for each PTM. A knowledge of these substrates will not only reveal how StUBK works, but will allow us also to fully understand the mode-of-action of PiSFI3 in targeting it to suppress early PTI transcriptional responses.

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Author contributions

PRJB, MJB, QH and ZT planned and designed the research. QH, HM, RKH, PCB, MA and YL, performed experiments and analysed data. QH, PRJB and MJB wrote the manuscript with input from all authors.

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**Supporting Information**

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Fig. S1 Representative leaf image showing *Phytophthora infestans* lesions following overexpression of each construct (GFP–PiSFI3, NLS/GFP–PiSFI3, or NES/GFP–PiSFI3) in *Nicotiana benthamiana*.

Fig. S2 PiSFI3 potato overexpression transgenic lines show enhanced *Phytophthora infestans* colonization.

Fig. S3 Transcript accumulation of flg22 maker genes posttreatment with flg22 on potato.

Fig. S4 SFI3 does not suppress programmed cell death triggered by INF1 or Cf4/Avr4.

Fig. S5 Alignment of UBK/PUB33.

Fig. S6 PiSFI3 interacts with a potato protein containing U-box and kinase domains.

Fig. S7 Silencing UBK by VIGS in *Nicotiana benthamiana* or by RNAi in potato..

Fig. S8 Silencing *SiUBK* by RNAi in potato enhance *Phytophthora infestans* colonization.

Fig. S9 VIGS of UBK does not compromise INF1 or Cf4/Avr4 cell deaths.
Fig. S10 Expression profile of *StUBK* posttreatment with flg22 and flg22 maker gene expression on *StUBK* RNAi lines post flg22 treatment.

Fig. S11 Overlays of the structures of the WY-domains of SFI3, Avr3a11 and PexRD2.

Fig. S12 SFI3 mutants lose nucleolar localisation.

Table S1 Primers used in this study.

Table S2 X-ray data collection and refinement statistics

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