

Fungal phytopathogens encode functional homologues of plant rapid alkalization factor (RALF) peptides

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SUMMARY

In this article, we describe the presence of genes encoding close homologues of an endogenous plant peptide, rapid alkalization factor (RALF), within the genomes of 26 species of phytopathogenic fungi. Members of the RALF family are key growth factors in plants, and the sequence of the RALF active region is well conserved between plant and fungal proteins. RALF1-like sequences were observed in most cases; however, RALF27-like sequences were present in the *Sphaerulina musiva* and *Septoria populicola* genomes. These two species are pathogens of poplar and, interestingly, the closest relative to their respective RALF genes is a poplar RALF27-like sequence. RALF peptides control cellular expansion during plant development, but were originally defined on the basis of their ability to induce rapid alkalization in tobacco cell cultures. To test whether the fungal RALF peptides were biologically active in plants, we synthesized RALF peptides corresponding to those encoded by two sequenced genomes of the tomato pathogen *Fusarium oxysporum* f. sp. *lycopersici*. One of these peptides inhibited the growth of tomato seedlings and elicited responses in tomato and *Nicotiana benthamiana* typical of endogenous plant RALF peptides (reactive oxygen species burst, induced alkalization and mitogen-activated protein kinase activation). Gene expression analysis confirmed that a RALF-encoding gene in *F. oxysporum* f. sp. *lycopersici* was expressed during infection on tomato. However, a subsequent reverse genetics approach revealed that the RALF peptide was not required by *F. oxysporum* f. sp. *lycopersici* for infection on tomato roots. This study has demonstrated the presence of functionally active RALF peptides encoded within phytopathogens that harbour an as yet undetermined role in plant–pathogen interactions.

Keywords: *Fusarium*, phytopathogen effectors, rapid alkalization factor (RALF).

INTRODUCTION

Fungal phytopathogens are a major threat to all plant ecosystems, both natural and agricultural (Stukenbrock and McDonald, 2008). Therefore, it is crucial to understand how fungal phytopathogens have co-evolved with their plant hosts to enable better management of crop losses and preserve natural biodiversity. The mechanisms by which pathogenic fungi manipulate their plant hosts are complex and diverse. It is now recognized that pathogen effector molecules (proteins or secondary metabolites) play a significant role in manipulating host plants to facilitate infection (de Jonge *et al.*, 2011). For example, biotrophic fungi produce effectors that allow the fungus to elude or manipulate host defences. Conversely, necrotrophic fungi produce effectors that induce host cell necrosis, thereby producing dead plant tissue for the fungus to colonize. Some effectors are vital for host infection; others are not strictly necessary, but assist in the full exploitation of the host resources (De Wit *et al.*, 2009). Most fungal effector genes remain undiscovered, but the increasing availability of fully sequenced fungal genomes has enabled new methods for the identification of putative effector genes using comparative genomics and candidate gene prediction tools.

The advent of next-generation sequencing (NGS) has dramatically increased the number of whole-genome sequences available for organisms from all kingdoms of life. Access to these data has allowed researchers to compare genes from disparate organisms (Thynne *et al.*, 2015b) and has provided significant insights into the genes involved in phytopathogen emergence and niche specialization (Gardiner *et al.*, 2012; Klosterman *et al.*, 2011), including the identification of pathogen genes related to host genes, either through horizontal gene transfer (HGT) or convergent evolution. By studying the molecular evolution of these genes, we can

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develop hypotheses about their function, including their potential roles as effectors.

Horizontal transfer of pathogenicity genes between fungi is a well-known phenomenon and an active area of research (Friesen *et al.*, 2006; Gardiner *et al.*, 2012; Klosterman *et al.*, 2011; Richards *et al.*, 2011; Thynne *et al.*, 2015b), although the exact mechanisms of genetic transfer are poorly understood. Recent studies have also demonstrated a role for plant to microorganism gene transfer leading to increased virulence. For example, the secreted proteinaceous *Verticillium* effector Ave1, which promotes pathogen virulence on the plant host, is hypothesized to have a plant origin (de Jonge *et al.*, 2012). Other examples exist in which plant-interacting organisms have apparently acquired plant growth and cell structure alteration factors. For example, C-terminally encoded peptides (CEPs) have been identified in the genome assemblies of phytopathogenic root knot nematodes (Bobay *et al.*, 2013). Similarly, homologues of plant expansins have been identified in the genomes of a range of bacteria and fungi (Nikolaidis *et al.*, 2014). CEPs and expansins are growth regulatory and plant cell-loosening molecules, respectively (Cosgrove, 2000; Delay *et al.*, 2013). These molecules have been postulated to assist in plant colonization, inhabitation and/or parasitization (Soanes and Richards, 2014). Pathogen genes of plant origin are therefore promising targets for further study as potential virulence factors.

The plant rapid alkalization factor (RALF) genes encode secreted peptides that were first identified through their ability to trigger a rapid increase in extracellular pH when added to plant cell suspensions (Pearce *et al.*, 2001). RALF peptides have roles in the development and regulation of plant roots, root hairs, legume nodules and pollen tubes (Murphy and De Smet, 2014; Pearce *et al.*, 2010). Subsequent *in silico* analyses of the *Arabidopsis thaliana* genome have identified 34 genes within the RALF family (Olsen *et al.*, 2002). These genes were named RALF1 to RALF34, and all homologues since identified in other species have been named according to their similarities to members of the *A. thaliana* family. Functional studies on a tobacco RALF peptide have revealed that proteolytic cleavage of a larger precursor protein generates a 49-amino-acid active peptide (Fig. 1) (Srivastava *et al.*, 2009). Mutational analysis has identified several motifs essential for RALF activity, including a YISY motif and four conserved cysteine residues that are required for disulphide bonding (Pearce *et al.*, 2010). Recently, the *Arabidopsis* receptor for RALF1 was identified as the plasma membrane receptor kinase FERONIA (FER). Activation of FER results in phosphorylation of plasma membrane H⁺-ATPase 2 at serine-899, causing inhibition of proton transport (Haruta *et al.*, 2014).

In this study, we describe the distribution of genes encoding RALF homologues amongst a diverse range of plant-pathogenic fungal species and RALF-like hybrid genes in a limited number of

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MDKSFTLFLTLTLVVFIISSPPVQAGFANDLGGVAVWATTGDNGSGCHGSAIECIGAEEM
↓
DSEINRRILATTKYISYQSLKRNSVPCSRRGASYNCQNGAQANPYSRGCSKIARCRS
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Fig. 1 Sequence of the *Nicotiana tabacum* rapid alkalization factor (RALF) peptide (NtRALF). The underlined region represents the predicted signal peptide and the arrow indicates the point of cleavage by a subtilisin-like serine protease (Srivastava *et al.*, 2009). The sequence in red represents the active RALF peptide. Asterisks (*) denote the essential YSIY motif. Cysteine residues required for disulphide bond formation and RALF peptide activity are denoted by hashtags (#) (Pearce *et al.*, 2010).

bacterial species. Functional testing of synthetic versions of two fungal RALF peptides showed that plants perceive and respond to the fungal peptides. These data suggest that these fungal RALF genes enable pathogenic fungi to manipulate host plant physiology.

RESULTS

Distribution and variation of RALF peptide sequences in the fungal and bacterial kingdoms

The search programs BLASTP and tBLASTn were used to screen publicly available fungal genomes for homologues of plant RALF proteins using the full-length protein sequences of *Arabidopsis thaliana* plant RALF genes 1–34 (Table 1). Using this approach, 26 different species of fungi were found to possess genes encoding the processed RALF peptide domain (Fig. 2, Table 1). These include the class Pucciniomycetes of the Basidiomycota and the classes Dothideomycetes and Sordariomycetes of the Ascomycota. All of the fungi found to possess RALF homologues are plant pathogens. Both biotrophic and necrotrophic fungi are represented in this list. Phylogenetic analysis of plant and fungal RALF peptide sequences showed that the fungal RALF homologues are interspersed amongst the plant RALFs (Fig. S1, see Supporting Information). Many of the fungal homologues resolved with the plant RALF1 proteins (Fig. S1). The *Pseudocercospora fijiensis* RALF homologue clustered with RALF homologues of *Musa acuminata malaccensis*. However, this was only supported with a low posterior probability of 33%. The *Septoria populicola* and *Sphaerulina musiva* RALF homologues clustered with the plant RALF27-like proteins with 100% posterior support (Figs 2 and S1). The best BLASTP hit to a complete plant protein for both fungal sequences was a RALF27-like protein from *Populus trichocarpa* (E-values of 3e-14 and 4e-13); however, this was only weakly supported by the gene tree analysis, with a posterior probability of 50% (Figs 2 and S1). It should be noted that there was no detectable homology between the N-terminal pro-peptide domains of the plant and fungal RALF sequences.

Analysis of the *Fusarium* species showed evidence of four divergent groups of RALF homologues (Fig. 3, Table 2), which were designated as Groups I–IV. Members of Group I are present

Table 1 Fungal species identified as containing rapid alkalization factor (RALF) homologues.

Fungi containing RALF sequences	JGI/NCBI/GenBank/Ensembl/UniProt sequence identifier
Ascomycetes	
Dothideomycetes	
Botryosphaeriaceae	
Botryosphaeriaceae spp.	See Table 3
Corynesporascaceae	
<i>Corynespora cassiicola</i>	jgi Corca1 631921
Leptosphaeriaceae	
<i>Leptosphaeria biglobosa</i>	GenBank: F0905662.1
<i>Leptosphaeria maculans</i>	NCBI Reference Sequence: XP_003844940.1
<i>Phoma tracheiphila</i>	jgi Photr1 406253
Mycosphaerellaceae	
<i>Pseudocercospora fijiensis</i>	NCBI Reference Sequence: XP_007921540.1
<i>Septoria populiicola</i>	jgi Seppo1 99826
<i>Sphaerulina musiva</i>	GenBank: EMF16709.1
Pleosporaceae	
<i>Pyrenophora tritici-repentis</i>	NCBI Reference Sequence: XP_001937123.1
<i>Pyrenophora teres</i>	jgi Pyrtt1 194499:17017-24160
<i>Setosphaeria turcica</i>	jgi Settu1 scafd_12:1053528-1053731
Teratosphaeriaceae	
<i>Teratosphaeria nubilosa</i>	jgi Ternu1 214030
Sordariomycetes	
Glomerellaceae	
<i>Colletotrichum higginsianum</i>	GenBank: CCF44719.1
Plectosphaerellaceae	
<i>Verticillium dahliae</i>	GenBank: CP009075.1
<i>Verticillium alfalfae</i>	
Nectriaceae	
<i>Fusarium</i> spp.	See Table 2
Basidiomycetes	
Pucciniomycetes	
Melampsoraceae	
<i>Melampsora lini</i>	jgi Melli1 sc_1450:29870-30703
<i>Melampsora larici-populina</i>	NCBI Reference Sequence: XP_007408986.1
Cronartiaceae	
<i>Cronartium quercuum</i>	jgi Croqu1 50147

in all isolates of *F. oxysporum* (*Fox*) that have been sequenced to date, although the Group I RALF genes of two *Fox* isolates encode sequence variations suggesting loss of function (a premature stop codon in the human pathology isolate FOSC 3-a, and a tyrosine in place of a conserved cysteine in the non-pathogen Fo47). Group I can be subdivided into two subgroups A and B, with subform IB also present in *F. verticillioides*, *F. fujikuroi* and *F. circinata* (Fig. 3). Interestingly, subform IA distinguishes two tomato pathogens, *F. oxysporum* f. sp. *lycopersici* (*Fol*) isolate MN25 and *F. oxysporum* f. sp. *radicis-lycopersici* (*Forl*) from a third *Fol* isolate 4287. The divergent *Fol* homologues, here designated RALF-B (from 4287) and RALF-C (from MN25), were chosen for further study.

The Group II family of RALF homologues has only two members, one from *Fox* FOSC 3-a, which could potentially complement the aberrant FOSC 3-a Group I homologue, and the other from *F. oxysporum* f. sp. *melonis*, which contains an apparent frameshift mutation (Fig. 3). The Group III family of RALF homologues has only three members, one from *Forl* CL57 and the others from *F. graminearum* and *F. pseudograminearum* (Fig. 3).

Genes encoding a fourth family of RALF-like proteins (Group IV) were also found in *Fusarium*. These differ from the three families described above by a central deletion encompassing the conserved cysteines that form the first disulphide bond (Fig. 3). Nevertheless, the sequence conservation within this family suggests the retention of function, but whether they possess the same function as the fungal RALF homologues described above requires further investigation. This gene family can also be divided into two sequence-divergent subgroups that seem to be distributed in a mutually exclusive fashion between different *formae speciales* of *Fox*, although two (*vasinfectum* and *Fol* 4287) do not possess either form. Interestingly, Group IV genes seem to be absent from plants and other fungal genomes, suggesting that they are unique to *Fusarium*.

Notably, all four possible combinations of the Group I and IV RALF variants occur in *Fox*, indicating a polyphyletic origin for at least some of these gene combinations. This is a curious finding given that these genes also occur in *F. verticillioides*, *F. fujikuroi* and *F. circinata*, suggesting that they are located on the core conserved set of non-mobile chromosomes in *Fusarium*, rather than the mobile lineage-specific chromosomes that seem to determine host specificity in *Fox* (Ma *et al.*, 2010). Indeed, the RALF-B gene is encoded by core chromosome 12 in *Fol* 4287.

Subsequent to the preliminary database search, we annotated a number of new RALF sequences within the genomes of members of the Botryosphaeriaceae. Similarly to the *Fusarium* spp., members of the Botryosphaeriaceae family showed evidence of their own three divergent groups of RALF homologues (Table 3, Fig. S2, see Supporting Information). These three groups were designated BI–BIII. Amino acid variations between these three divergent groups are highlighted in Fig. S2A. BI contains an amino acid string ('ISNGAM') apparently rare among both fungal RALFs (only non-Botryosphaeriaceae RALF is in *Colletotrichum higginsianum*) and plant RALFs (only observed in *Brachypodium distachyon*). BII is similar in amino acid composition to the *Fusarium* Group I. BIII is similar, and perhaps divergent from, BII. *Botryosphaeria dothidea* contains two RALF homologues (BI and BII). *Botryosphaeria dothidea*'s BI RALF homologue was previously unannotated. *Diplodia seriata* contains one RALF homologue (BIII). *Neofusicoccum parvum* contains one RALF homologue (BI). *Macrophomina phaseolina* contains four RALF homologues (BI, two BIIIs and BIII). All but one of *M. phaseolina*'s RALFs were previously un-annotated. With four predicted RALF peptides,

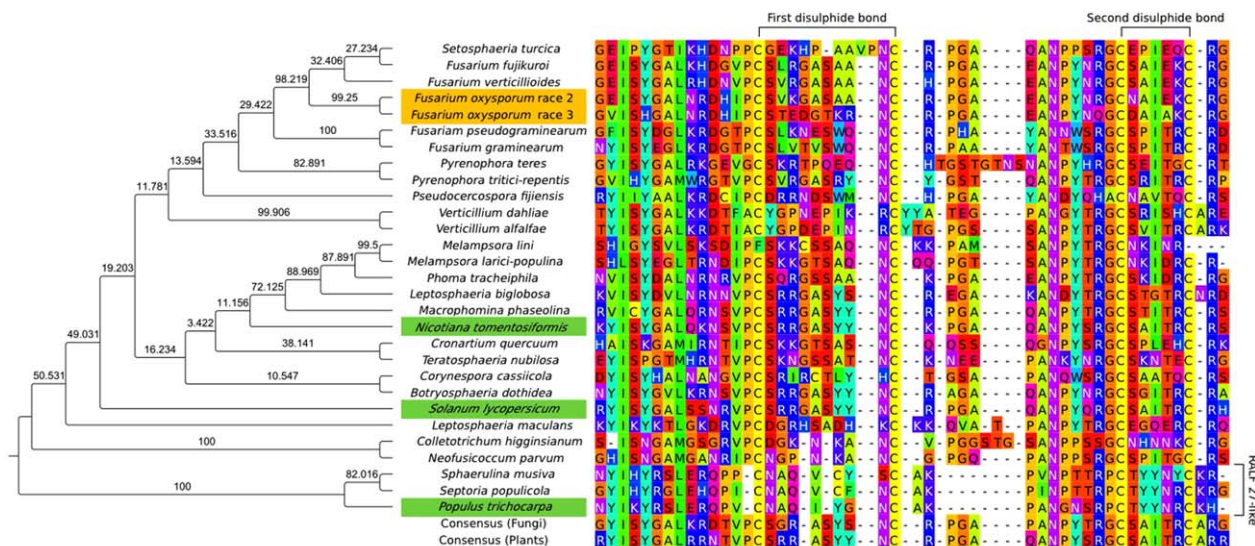


Fig. 2 Identification, alignment and analysis of the rapid alkalization factor (RALF) domain in fungi and selected plants. Aligned RALF peptide domain sequences from five isolates of *Fusarium oxysporum*, 24 other fungal species and selected plants (highlighted in green) are presented. Cysteine pairs expected to form disulphide bonds are labelled at the top. The *Melampsora lini* RALF may have an alternative disulphide bond, marked with a dotted line. Support values on each branch refer to the percentage posterior probability of each node. RALF sequences from two species of fungi, *Septoria populicola* and *Sphaerulina musiva*, clustered with plant RALF27-like genes with 100% posterior support (labelled on the right). The plant RALF27-like sequences are from *Populus trichocarpa* (gi:566213100) and *Arabidopsis thaliana* (gi:15230083). The other plant RALF sequences are from *Solanum lycopersicum* (gi:460366641), corresponding to the peptide used here in seedling and cell culture assays, and from *Nicotiana tomentosiformis* (gi:697149758), which is identical to the original RALF sequence reported from *N. tabacum* and *A. thaliana* RALF1 (gi:15218637). The divergent RALF sequences of *Fusarium oxysporum* isolates 4287 and MN25 are highlighted in gold. Individual amino acids are coloured using the Taylor scheme (Taylor, 1997).

M. phaseolina has the highest number of predicted fungal RALF peptides identified in this study. One of the BII RALF peptides shares a conserved RALF peptide domain with the mature RALF peptide from soybean species (*Glycine max* and *Glycine soja*). For this region, these species have a 100% amino acid identity. Variation among these species is found in the N-terminal and C-terminal regions. In addition, common bean (*Phaseolus vulgaris*) has a 98% sequence similarity for this conserved region, and mung bean (*Vigna radiata*) has a 100% sequence similarity for 97% of the query coverage. The only species of Botryosphaeriaceae analysed in this study not to contain RALF peptides are three closely related species: *Eutiarospora darliae*, *E. pseudodarliae* and *E. tritici-australis*. These are the causal agents of white grain disorder (WGD) in wheat (Thynne *et al.*, 2015a).

We re-performed the database searches using BLASTp, this time searching the bacterial portal of the National Center for Biotechnology Information (NCBI) database. We identified certain members of *Actinobacteria* that possess putative secreted proteins with an incorporated RALF peptide domain motif (Fig. S3, see Supporting Information). In both fungal and plant RALFs, this particular motif is located at the C-terminus of the mature peptide. Similarly, in these bacterial proteins, the RALF domain motif is at the C-terminus of the protein. Unlike plant and fungal RALFs, the bacterial RALF domain-containing proteins also contain a domain

homologous to the S1 pertussis toxin subunit. These bacterial species include a number of plant-pathogenic species, e.g. *Streptomyces acidiscabies* (a pathogen of potatoes) (Huguët-Tapia *et al.*, 2012).

Tomato seedling growth and development are severely inhibited by a synthetic RALF-B peptide

Endogenous RALF peptides arrest the growth and development of plant roots (Pearce *et al.*, 2010). To determine whether fungal RALF peptides share this property, we synthesized RALF peptides based on RALF2 of tomato (SI-RALF, NCBI GenBank GI:460366641) and those of *Fol* race 2 isolate 4287 and race 3 isolate MN25 (RALF-B and RALF-C, respectively) (Figs 2 and S4, see Supporting Information). *Fol* 4287 was the first fully sequenced *Fox* isolate, and MN25 is a representative of race 3, which appeared in the 1980s and overcame resistance to races 1 and 2 (Bournival *et al.*, 1989). A non-functional mutant of SI-RALF lacking the required YISY motif (SI-RALF Δ) was used as a negative control. Germinating tomato seeds were grown in medium containing each RALF peptide at 10 μ M (Fig. 4), or with no added peptide. Untreated seedlings remained healthy and showed normal development of roots and shoots. Seedlings grown in medium containing SI-RALF developed less biomass than the untreated controls, but this was not significantly different from the SI-RALF Δ

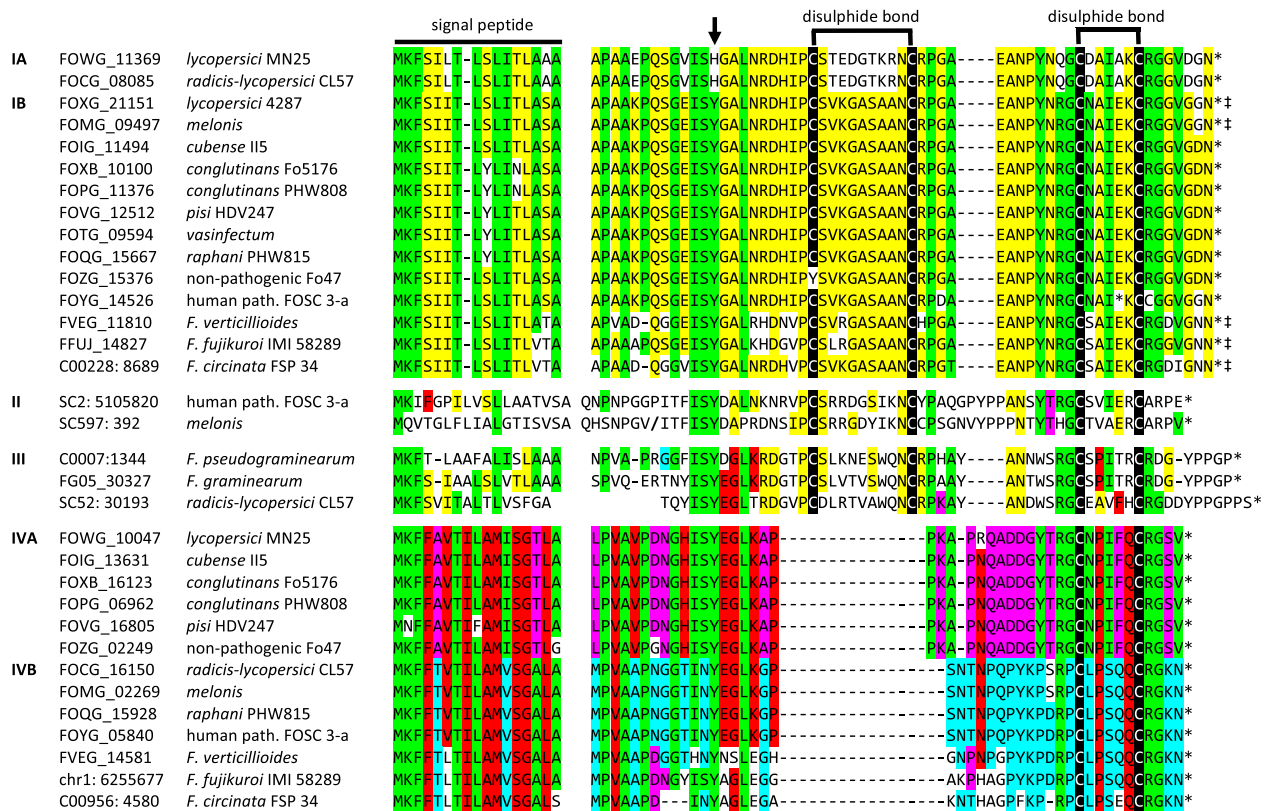


Fig. 3 Multiple sequence alignments of *Fusarium* rapid alkalization factor (RALF) and RALF-like homologues. Group I comprises a large family of RALF homologues from *Fusarium verticillioides*, *F. fujikuroi*, *F. circinata* and various *formae speciales* of *F. oxysporum* (*Fox*). It should be noted that the Group IA RALF homologues FOWG_113369 *F. oxysporum* f. sp. *lycopersici* 4287 and FOCG_08085 *F. oxysporum* f. sp. *radicis-lycopersici* CL57 are very similar to one another, but divergent from the Group IB RALF sequences at specific points within the predicted mature RALF peptide, notably the sequence between the first two cysteines and a histidine substitution in place of a conserved tyrosine (arrowed). Four cysteine residues involved in disulphide bonding are conserved in all isolates, except FOZG_15376, where the first cysteine is swapped at position 39 to a tyrosine. FOYG_14526 has an early stop codon at site 67, prior to the final cysteine at site 69. Group II comprises a small family of divergent RALF homologues from *F. oxysporum* f. sp. *melonis* and a human pathology isolate of *Fox*. It should be noted that the f. sp. *melonis* homologue has an apparent frameshift indicated by a slash (/) that could be the consequence of a sequencing error rather than a frameshift mutation. Group III comprises a small family of divergent RALF homologues from *F. oxysporum* f. sp. *radicis-lycopersici* CL57, *F. graminearum* and *F. pseudograminearum*. Group IV comprises a large family of RALF homologues that have an internal deletion relative to the RALF homologues shown in Groups I, II and III. This deletion includes the cysteine residues that form the first disulphide bond indicated above the alignments. It should be noted that this family contains two apparently mutually exclusive subforms, as can be seen from the conserved residues highlighted in pink and blue. Conserved cysteines are highlighted in black. Other residues conserved between Groups I and IV, and, to some extent, between Groups II and III are highlighted in green. Residues conserved within Group I and, to some extent, Groups II and III are highlighted in yellow. Residues conserved within Group IV and, to some extent, Groups II and III are highlighted in red. Residues conserved within Group IV, but differing between Groups IVA and IVB, are highlighted in pink and blue. Signal peptide cleavage sites were predicted using SignalP 4.1 (<http://www.cbs.dtu.dk/services/SignalP/>). Disulphide bond locations are based on Pearce *et al.* (2001). †Transcript and protein sequences predicted using the same splice acceptor site as predicted for all of the other sequences in this group. *Stop codon.

treatment (Fig. 4A). However, the SI-RALF-treated seedlings were developmentally stunted with minimal root formation compared with the SI-RALFA-treated seedlings (Fig. 4B). The growth and root development of seedlings treated with synthetic RALF-B peptide were severely inhibited relative to mock- and SI-RALFA-treated seedlings. This was reflected in their significantly lower average weights. In contrast, the synthetic RALF-C peptide did not inhibit root growth. There was no significant difference in average biomass between seedlings treated with RALF-C and the mock and SI-RALFA treatments. Similarly, the seedlings appeared healthy with well-developed root systems.

The RALF-B peptide elicits recognized plant RALF responses

The receptor for the *Arabidopsis* RALF1 peptide is the CrRLK1L family receptor FER (Haruta *et al.*, 2014). As FER function is associated with the accumulation of extracellular reactive oxygen species (ROS) (Wolf and Höfte, 2014), and this is also a typical consequence of plant defence activation, we asked whether synthetic *Fol* RALF peptides could also elicit ROS production. Treatment of leaf discs of tomato and another solanaceous species, *Nicotiana benthamiana*, with SI-RALF at 10 μM induced a ROS

Table 2 *Fusarium* isolates and their rapid alkalization factor (RALF) homologue IDs or DNA sequence origins.

Fusarium isolate	Protein ID/origin
<i>Fusarium fujikuroi</i> IMI 58289	FFUJ_14827 chromosome 1: 6255677–6255856
<i>Fusarium graminearum</i> PH-1 (NRRL 31084)	FG05_30327
<i>Fusarium oxysporum</i> f. sp. <i>conglutinans</i> PHW808 (NRRL 54008)	FOPG_11376 FOPG_06962
<i>Fusarium oxysporum</i> f. sp. <i>conglutinans</i> Fo5176	FOXB_10100 FOXB_16123
<i>Fusarium oxysporum</i> f. sp. <i>cubense</i> TR4 II5 (NRRL 54006)	FOIG_11494 FOIG_13631
<i>Fusarium oxysporum</i> f. sp. <i>lycopersici</i> 4287 (NRRL 34936)	FOXG_21151
<i>Fusarium oxysporum</i> f. sp. <i>lycopersici</i> MN25 (NRRL 54003)	FOWG_11369 FOWG_10047
<i>Fusarium oxysporum</i> f. sp. <i>melonis</i> (NRRL 26406)	FOMG_09497 FOMG_02269
<i>Fusarium oxysporum</i> f. sp. <i>pisi</i> HDV247 (NRRL 37622)	Super Contig 597: 392–624 FOVG_12512 FOVG_16805
<i>Fusarium oxysporum</i> f. sp. <i>radicis-lycopersici</i> CL57 (NRRL 26381)	FOCG_08085 FOCG_16150
<i>Fusarium oxysporum</i> f. sp. <i>raphani</i> PHW815 (NRRL 54005)	Super Contig 52: 30193–30411 FOQG_15667 FOQG_15928
<i>Fusarium oxysporum</i> f. sp. <i>vasinfectum</i> (NRRL 25433)	FOTG_09594
<i>Fusarium oxysporum</i> human path. isolate FOSC 3-a (NRRL 32931)	FOYG_14526 FOYG_05840
<i>Fusarium oxysporum</i> non-pathogenic Fo47 (NRRL 54002)	FOZG_15376 FOZG_02249
<i>Fusarium pseudograminearum</i> CS3096	Contig 007: 1344–1371
<i>Fusarium verticillioides</i> 7600 (NRRL 20956)	FVEG_11810 FVEG_14581
<i>Fusarium circinata</i> FSP34	Contig 00228: 8689–8910 Contig 00956: 4580–4747

burst. The ROS burst elicited by SI-RALF was stronger in tomato than in *N. benthamiana*. In contrast, RALF-B (but not RALF-C) induced a massive ROS burst in both solanaceous species, only minutes after treatment. RALF-C was not active in these assays. Neither the mock nor SI-RALF Δ treatments elicited a significant ROS response in the leaves of either species.

The dose response for fungal RALF peptides was also determined. As the ROS bursts induced by RALF-B were comparable between tomato and *N. benthamiana*, the activity of this peptide was tested in *N. benthamiana*. RALF-B was active down to a concentration of 10 nM, with progressive increases in ROS output observed up to 500 nM (Fig. S5, see Supporting Information). In

Table 3 Botryosphaeriaceae isolates and their rapid alkalization factor (RALF) homologue IDs or DNA sequence origins.

Botryosphaeriaceae isolate	Protein ID/origin
<i>Botryosphaeria dothidea</i>	gjilBotdo1_1 289944 Node_8012: (92987–93247)
<i>Diplodia seriata</i>	gjil821070376
<i>Macrophomina phaseolina</i>	gjil407928039 Contig_00502: (91876–92412) Contig_00151: (16464–16099)
<i>Neofusicoccum parvum</i>	Contig_00254: (124817–124602) gjil615408450

contrast, ROS output subsequent to increasing concentrations of RALF-C did not exceed control levels, providing further evidence that the peptide does not elicit a typical RALF response in plants.

The ability of RALF-B to induce alkalization was also determined. RALF peptides were originally named as such because of their strong and rapid alkalization of tobacco suspension-cultured cells (Pearce *et al.*, 2001). To test the ability of RALF-B to induce alkalization, the peptide was incubated with cut leaf strips from both tomato and *N. benthamiana*. For both plants, RALF-B elicited an alkalization response not significantly different from that of the tomato RALF peptide (Fig. 5C,D).

Similarly, the activation of mitogen-activated protein kinases (MAPKs), a known RALF response (Pearce *et al.*, 2001), was also investigated. Peptide-induced MAPK activation was measured in *N. benthamiana* leaf discs as confirmation of the variation of activity in response to the different peptides. Treatments with 10 μ M SI-RALF or RALF-C activated MAPK progressively, with greater activity detected at 15 min after peptide addition (Fig. S6, see Supporting Information). Interestingly, the addition of only 100 nM RALF-B (i.e. 100 times less than that of SI-RALF) caused strong activation of MAPK that peaked at 5 min and showed a decline by 15 min. This mirrors the ROS profile arising from RALF-B treatment. Overall, these data (ROS burst, alkalization and MAPK activation) prove that plant cells perceive fungal RALF peptides.

The *Fol* RALF-B gene is expressed during infection of tomato roots

If RALF-B plays a role in fungal pathogenicity, it should be expressed during infection of tomato roots by *Fol* isolates carrying the RALF-B gene. A reverse transcriptase-polymerase chain reaction (RT-PCR) analysis was carried out on roots of the susceptible tomato cultivar M82 infected with *Fol* isolate #1943, which carries the RALF-B gene, at 3 and 6 days post-inoculation (dpi). No RALF-B expression was detected in samples from the roots of mock-inoculated tomato plants or 5-day-old mycelia grown *in vitro*, but

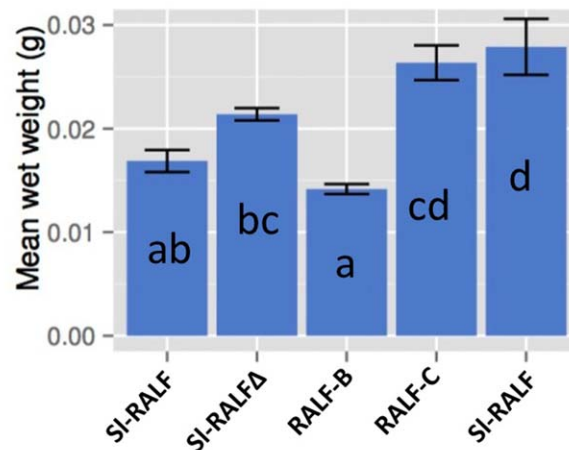
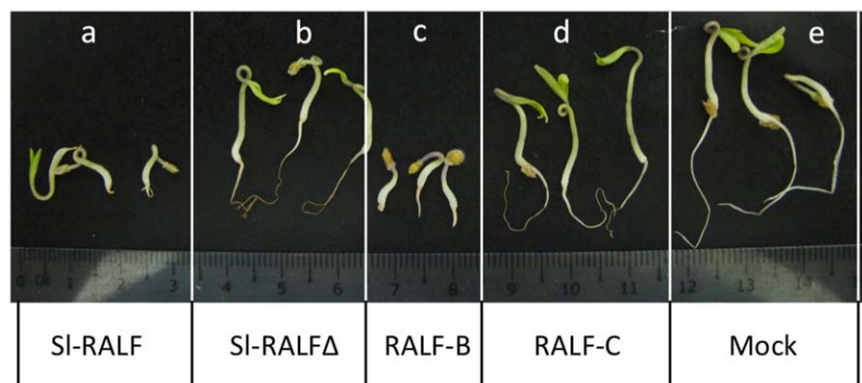


Fig. 4 Effect of rapid alkalization factor (RALF) peptides on germinating tomato seedlings. (A) Mean weights were calculated from whole seedlings grown in medium containing 10 μ M synthetic RALF peptides ($n = 9$). The letter codes a–d indicate statistical significance: treatments which do not share any letter codes are significantly different ($P < 0.05$, corrected for multiple testing). Error bars correspond to standard error. (B) Digital images of representative samples in (A): a, SI-RALF; b, SI-RALF Δ ; c, RALF-B; d, RALF-C; e, mock treatment.



expression was detected in the 3- and 6-dpi samples (Fig. 6). The *FEM1* gene (Schoffemeer *et al.*, 2001) was used as a positive control for fungal gene expression. The RALF-B primers flanked an intron, thus yielding different-sized products for cDNA and genomic DNA (Fig. 6). No RALF-B or *FEM1* genomic DNA products were detected in cDNA samples (Fig. 6), and no RALF-B or *FEM1* products were detected in samples prepared without reverse transcriptase (not shown), indicating that the RALF-B and *FEM1* bands observed at 3 and 6 dpi are authentic RT-PCR products. The absence of RALF-B expression in mycelia grown *in vitro* suggests that RALF-B expression is induced during plant infection, consistent with a potential role as an effector.

Effect of knocking out the RALF-B gene in *Fol* race 3

To test the role of the RALF-B gene in *Fol* pathogenicity, knockout (Δ RALF) mutants were obtained by *Agrobacterium*-mediated transformation, using a binary T-DNA vector carrying a hygromycin resistance gene as a selectable marker for gene replacement and a thymidine kinase gene as a counter-selectable marker against ectopic T-DNA insertions (Fig. S7A, see Supporting Information). A total of 44 *Fol* transformants that showed resistance to hygromycin and 5'-fluoro-2'-deoxyuridine were screened by PCR,

confirming deletion of the RALF coding sequence in four transformants (named $\Delta 2$, $\Delta 23$, $\Delta 24$ and $\Delta 31$) (Fig. S7B).

The four *Fol* Δ RALF transformants were used to evaluate the effect of the loss of the RALF gene on virulence. The roots of 3-week-old plants of the tomato cultivar M82 (susceptible to *Fol* race 3) were wounded and subsequently inoculated with *Fol* race 3, a transformant with an ectopic insertion of the T-DNA (i.e. no deletion of RALF gene) and the two Δ RALF transformants. The disease symptoms were recorded and disease scores were assigned as described by Rep *et al.* (2004), with scores of 0 = healthy plant to 4 = severely wilted plant.

No consistent significant differences in fungal virulence were found between wild-type *Fol* race 3 and any of the transformants (Fig. S7D,E). Δ RALF transformants not the ectopic mutant strain tested. Therefore, using a wound infection assay, we found no strong evidence that the RALF-B gene is involved in the pathogenicity of *Fol* race 3 on tomato.

DISCUSSION

Here, we have reported that a number of fungal plant pathogens encode RALF-like peptides in their genomes, and have provided evidence that at least a subset of these peptides can be perceived by plant cells. RALF peptides have high levels of conservation

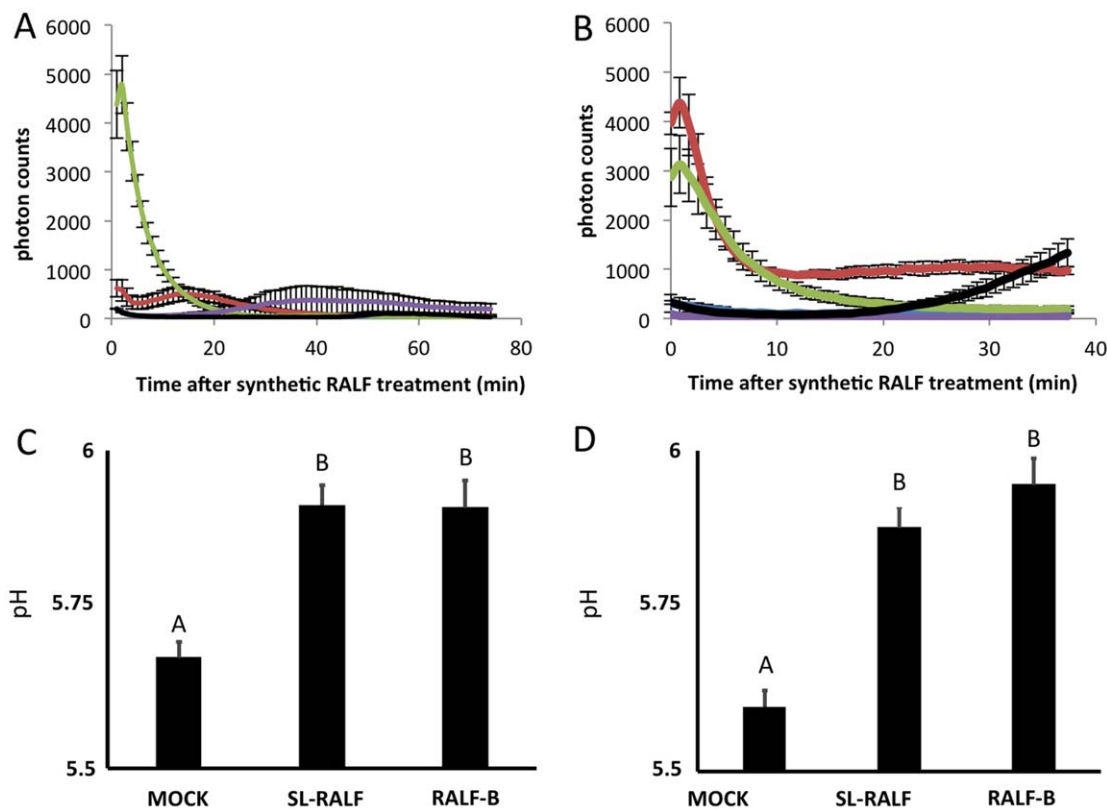


Fig. 5 Synthetic rapid alkalization factor (RALF) peptides induce reactive oxygen species (ROS) production and alkalization in *Nicotiana benthamiana* and tomato leaves. The ROS burst over time after treatment with 10 μM of each synthetic RALF peptide in *N. benthamiana* (A) and tomato (B) leaves. (C, D) Induction of alkalization in *N. benthamiana* and tomato, respectively, by RALF-B and SI-RALF. For (A) and (B), line colours are as follows: dark blue, mock treatment; red, SI-RALF; green, RALF-B; purple, RALF-C; black, SI-RALF Δ .

across a wide range of plant species, and the importance of their roles in the regulation of plant physiology is becoming increasingly apparent (Covey *et al.*, 2010; Murphy and De Smet, 2014; Pearce *et al.*, 2001). The suggestion that fungi deploy RALF peptides as effectors with the assumed consequence of improving fungal interaction with their hosts provides new perspectives both to the understanding of the pathogenic strategies of fungi and the evolutionary mechanisms underlying the acquisition of virulence.

Of the fungal genera carrying RALF homologues, the genus *Fusarium* has the most diverse array with four groups of RALF homologues (Fig. 3), with two subtypes in Groups I and IV. Each of the four groups has representatives in *Fox*, making *Fox* one of the most interesting fungal species for further study of fungal RALF peptides. However, it is possible that this apparent diversity is a result, in part, of the large number of *Fusarium* species, and *Fox formae speciales* in particular, whose genomes are publicly available. In bacteria, large numbers of genome sequences for a single species have catalysed the concept of the pan-genome. The pan-genome represents the entire genomic coding repertoire of all strains within a species, enabling intraspecific variation to be observed and assessed (Vernikos *et al.*, 2015). This variation can

be used to assist in explanations of varied lifestyles between isolates. The idea of the pan-genome is becoming more common in fungi and the re-sequencing of fungal genomes has begun to reveal the level of intraspecific variation (Thynne *et al.*, 2015b). With many re-sequenced *Fox* genomes and diversity in *Fox* lifestyles (non-pathogenic, plant-pathogenic, insect-pathogenic and opportunistic human pathogen), the genetic determinants underpinning this diversity can be identified and analysed. Focusing on the RALF sequences identified here, considerable intraspecific variation was observed between different *Fox* genomes. Intriguingly, the two sequenced *Fox* isolates that are not reported plant pathogens (isolate FOSC 3-a, which is a human clinical isolate, and isolate Fo47, which is non-pathogenic) carry mutations in conserved cysteine residues that should render the RALF peptide biologically inert. Disulphide bonds formed between the two pairs of cysteines in RALF are essential for activity (Pearce *et al.*, 2010). The RALF genes from these isolates appear to be undergoing pseudogenization, perhaps because they are not required outside of the plant-pathogenic niche. In this study, the *FoI* RALF-B variant elicited a strong response in the solanaceous plants tested. In contrast, there was a comparatively negligible response to *FoI* RALF-C, with

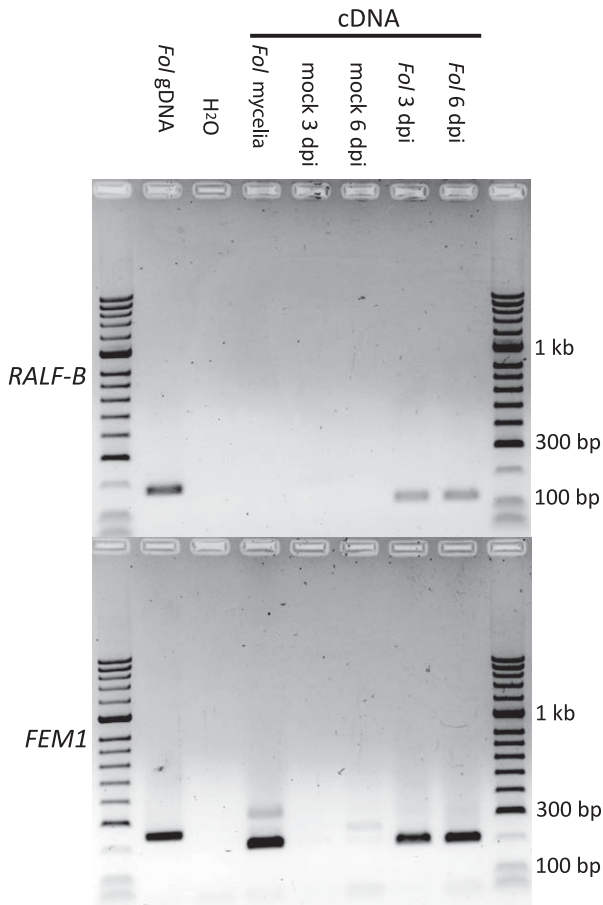


Fig. 6 Reverse transcriptase-polymerase chain reaction (RT-PCR) analysis shows that RALF-B is expressed during infection of tomato roots by *Fusarium oxysporum* f. sp. *lycopersici* (*Fol*). RT-PCR analysis of RALF-B (top gel image) shows bands (expected size, 123 bp) consistent with gene expression in *Fol*-infected roots at 3 and 6 days post-inoculation (dpi), but not in mock-inoculated tomato plants or 5-day-old mycelia grown *in vitro*. RT-PCR analysis of *FEM1* (bottom gel image) shows bands (expected size, 201 bp) consistent with expression in infected roots at 3 and 6 dpi and in 5-day-old mycelia grown *in vitro*. The *FEM1* control shows that cDNA synthesis was successful for the mycelial RNA sample and that the resulting cDNA sample supported PCR. PCR analysis of a *Fol* genomic DNA sample allowed the detection of the RALF-B and *FEM1* genes in *Fol*. The RALF-B and *FEM1* genomic DNA and cDNA products differ in size (expected sizes: 179 bp versus 123 bp for RALF-B and 250 bp versus 201 bp for *FEM1*), allowing the detection of genomic DNA contamination in cDNA samples. No genomic RALF-B or *FEM1* products were detected in the cDNA samples.

only weak activation of MAPK observed in treated *N. benthamiana*. Perhaps these differences are related to the swap of a conserved tyrosine with histidine early in the active peptide domain, as this tyrosine is important for RALF activity (Pearce *et al.*, 2010). RALF-C is found in *Fo1*, which causes crown rot of tomato, and therefore has a different pathogenic lifestyle to *Fol*. It is possible that the RALF-C variant plays a different role in *Fo1* than does RALF-B in *Fol*, but why then does *Fol* MN25 carry RALF-C instead

of RALF-B like *Fol* 4287? It is possible that *Fol* MN25 carries other genetic factors, such as the Group IV RALF-like genes present in *Fol* MN25, but absent from *Fol* 4287 (Fig. 3), which compensate for the absence of RALF-B. These are questions for further study.

Although the RALF variation found within *Fox* does not reveal the origins of the RALF gene family in fungi, their polyphyletic nature does suggest genetic mobility within *Fox*. All four pairwise combinations of the two apparently mutually exclusive subtypes of Group I and Group IV RALF homologues (Fig. 3) were found in different *formae speciales* of *Fox*, suggesting that one or both of these genes may be mobile, i.e. transmitted horizontally. This opens up yet another area for further investigation.

Using a synthetic RALF-B peptide, we demonstrated that a fungal RALF peptide can be perceived by plants. This is a highly significant result because it suggests that pathogenic fungi hijack endogenous plant physiological mechanisms to enhance their virulence. RALF was initially defined as a plant peptide able to trigger the alkalization of plant cell cultures, and subsequently demonstrated to play varied roles in plant development. (Murphy and De Smet, 2014; Pearce *et al.*, 2001). Here, we showed that treatment of plants with both plant and fungal RALF caused seedling growth inhibition, a rapid burst of ROS, MAPK activation and alkalization, all hallmarks of defence activation by plants. Elicitation of ROS by RALFs has not been reported previously and represents an important addition to the spectrum of responses associated with RALF perception. We tested two representative divergent RALF sequences from the 12 *Fox* sequences. Only one of these caused consistent activation of plant responses at very low concentrations. Interestingly, RALF-B was very much more potent than the endogenous SI-RALF peptide in treated *N. benthamiana*, and showed marked differences in both the extent and timing of the plant response. It is interesting to note that SI-RALF only induced a strong response in tomato, whereas RALF-B was able to induce a strong response in both tomato and *N. benthamiana*. This is consistent with the apparent conservation of RALF peptides across *Fox* spp. In these species, these peptides are located on the conserved (not lineage-specific) chromosomes. As such, promiscuity of activity among different potential hosts may be beneficial. At this stage, it is difficult to interpret the differences in activities between peptides, other than to point out that they must be a result of the sequence differences between the peptides. A key objective of this analysis was to demonstrate that a plant could perceive a fungal RALF peptide. This was achieved.

The expression of RALF-B in *Fol* during *in planta* growth was consistent with a potential role in mediating disease. However, under the conditions used for pathogenicity assays in this study, strains of *Fol* lacking RALF-B displayed similar disease symptoms to wild-type *Fol*, implying that the peptide is not involved in the induction of disease symptoms. These data suggest that RALF-B plays other as yet undetermined roles in this host–pathogen

interaction not trialled in this study. Inoculation was performed with cut roots, dipped in spore solution. This is a routine technique for the assay of *Fox* infections of tomato. Perhaps the fungal RALFs are involved in the manipulation of the host prior to and/or during colonization. Potential roles for RALF are the focus of ongoing research.

Whilst this article was still under review, Masachis *et al.* (2016) reported a functional role for RALF-B in *Fol* pathogenicity, but clear differences in the infection assays and scoring criteria were apparent when compared with this study. Masachis *et al.* (2016) grew plants for inoculation in vermiculite with no plant nutrients provided, and it is possible that assays conducted on such plants may have revealed an effect that was not evident in healthy plants grown in potting mix well supplied with nutrients. They also scored plant survival as opposed to disease symptoms and over a much longer period (up to 35 days as opposed to 21 days in our study), and it is interesting to note that their study also showed no significant effects at 21 dpi, i.e. no significant differences in survival. Therefore, we are confident that, under the inoculation conditions and scoring criteria used in this study, we found no significant effect of the RALF-B knockout on pathogenicity, and can only conclude that any effect of RALF-B on pathogenicity was too subtle for us to detect under our experimental conditions.

An intriguing observation from the distribution of RALFs among fungi was that the two poplar (*P. trichocarpa*) pathogens, *Sp. musiva* (*Mycosphaerella populorum*) and *Se. populicola* (*Mycosphaerella populicola*), encode RALF27-like homologues. Recent comparisons between the genomes of these two species have uncovered an adaptation event that probably altered the fitness of *Sp. musiva* to become more pathogenic to poplars in relation to *Se. populicola* (Dhillon *et al.*, 2015). Similarly, the gain or evolution of this peptide probably assisted the interaction of these fungi with poplars. Unlike most of the other fungal RALF homologues identified, the most similar homologues to the *Sp. musiva* and *Se. populicola* RALF27-like peptides are those of the host they infect. This implies that the fungal peptides have the same role in plants as the endogenous peptides. This is also a potentially interesting model for the study of the function of RALF27. No function has yet been assigned to this member of the RALF family, which is only briefly mentioned in the literature (Lafleur *et al.*, 2015; Marmiroli and Maestri, 2014; Olsen *et al.*, 2002). The characterization of the role of fungal RALF27 homologues during infection may also help to elucidate the function of the host homologue.

Similar to *Sp. musiva* and *Se. populicola*, both *Pseudocercospora fijiensis* and *M. phaseolina* share RALF homologues similar to their respective hosts. *Pseudocercospora fijiensis* has a RALF that resolves with banana (*Musa acuminata malaccensis*), and *M. phaseolina* has a RALF with sequence similarity to a range of bean species. The

RALF peptide from *M. phaseolina* is particularly interesting as it shares extremely high levels of sequence conservation for the peptide domain with a number of bean species (some of its primary hosts). However, this sequence clearly relates to other fungal RALF peptides, falling with a group of peptides shared among a range of fungal species (e.g. other members of the family Botryosphaeriaceae and the genus *Fusarium*). This would suggest that the peptide domain of this protein in *M. phaseolina* has probably evolved to be better attuned with that of its hosts.

The distribution of RALFs in the fungal kingdom is sporadic, with members represented from the Dothideomycete and Sordariomycete classes, as well as in the evolutionarily more distant Pucciniomycetes. Strikingly, the represented species are all economically significant phytopathogens. The strong conservation of the RALF peptide domain, and the *in planta* transcription and host perception of one such variant, represents important evidence that fungi have co-opted this endogenous host signalling system to enhance their fitness on plants. The question remains, however, what is the ancestral origin of these conserved genes? The data presented here are consistent with conflicting evolutionary scenarios; HGT or convergent evolution. At this time, we cannot comment with any certainty on which scenario is the most likely.

Although the primary focus of this analysis has been to present the findings that a number of members of the fungal kingdom possess RALF peptides, we also identified a limited number of plant-pathogenic bacteria which possess genes encoding amino acid sequences with an incorporated RALF motif domain. However, we did not explore any potential functional role for these proteins because of their striking differences from fungal and plant RALFs. The mature peptides of the RALFs discussed in fungi are homologous to the endogenous plant peptides. In contrast, the amino acid sequences in bacteria with a RALF-like domain appear to be an amalgamation of a C-terminal RALF domain and the S1 subunit of the pertussis toxin. Pertussis toxin has been studied in detail, particularly its role as a virulence factor in bacterial–animal pathosystems. However, how these two unrelated domains may perform in bacterial–plant interactions is currently unknown.

There remains much to learn about the function of RALF peptides and their role in plant growth and development. Adding to this mystery is how particular plant-pathogenic organisms, both fungal and bacterial, utilize variants of these peptides to facilitate host infection and/or disease. In this study, we have demonstrated that genomes of specific fungal and bacterial phytopathogens encode RALF peptides, at least some of which are functionally active. We have focused on the interaction between *Fol*'s expression of a particular RALF peptide and infection in tomato. However, there are a number of different RALF homologues utilized in various microorganism–plant pathosystems. We anticipate that this discovery will form the basis of an understanding of how

different RALF peptides are utilized by microorganisms to co-opt plant machinery to facilitate *in planta* growth.

EXPERIMENTAL PROCEDURES

Identification of RALF homologues in fungal sequence databases

Homologues of RALF were identified with BLAST searches against the fungal sequence databases in NCBI (<http://www.ncbi.nlm.nih.gov>) and available via the MycoCosm portal in JGI (<http://genome.jgi-psf.org/programs/fungi/index.jsf>). The queries used were the *Arabidopsis thaliana* RALF1–RALF34 protein sequences from NCBI (Table S1). Because of the variability in the N-terminal and C-terminal regions, the BLAST search queries were refined where needed to include only the mature RALF peptide region (Pearce *et al.*, 2010). BLASTP searches were also employed to find RALF homologues using fungal proteins with RALF annotations (Altschul *et al.*, 1990). No e-value cut-off was used. Instead, individual BLAST hits were assessed for the presence of RALF-associated domains (Murphy and De Smet, 2014), regardless of e-value. tBLASTn searches were used to find mRNA sequences and un-annotated or differently annotated RALF homologues on fungal genome contigs/chromosomes (Condon *et al.*, 2013; Gardiner *et al.*, 2012; Nemri *et al.*, 2014; Ohm *et al.*, 2012). Where annotations were not present or were different, the online Softberry server (<http://linux1.softberry.com/berry.phtml>) was used to predict mRNA and protein sequences employing the FGNEISH+ prediction algorithm (Solovyev *et al.*, 2006). Fungal RALF protein sequences were used as queries against the plant protein database in NCBI to find the top plant BLAST hits (Altschul *et al.*, 1990).

Alignment and analysis of RALF protein sequences

A representative database of plant RALF proteins was compiled by searching for any plant amino acid sequence in the RefSeq database (Pruitt *et al.*, 2007) labelled with the term 'RALF' or 'Rapid Alkalinization Factor'. To compare RALF protein sequences from fungi and plants, GLAM2 (Frith *et al.*, 2008) was used to identify the conserved region corresponding to the RALF domain from the plant sequences, conserved sites within the RALF domain and to align the conserved sites. Its sister program GLAM2S-CAN was then used to align conserved sites from the fungal sequences, which were then added to the plant alignment.

The Bayesian phylogenetics software ExaBayes (Aberer *et al.*, 2014) was then used to infer the posterior distribution of a RALF gene tree based on the aligned sequences. The priors for this analysis were the LG substitution matrix (Le and Gascuel, 2008) (used as a fixed amino acid model prior), and a uniform prior on tree topologies. Eight independent chains were run with different seeds, sampled once every 3000 states. Using Tracer v1.6 (<http://beast.bio.ed.ac.uk/Tracer>), five chains were observed to converge at the same likelihood after 1.8 million states, and so sampled trees and statistics from the subsequent 3.6 million states of those converged chains were concatenated. The estimated sample size of the concatenated log likelihood was 747, which suggests sufficient sampling of the posterior, as this is much greater than the minimum threshold of 200 (Kuhner, 2009).

A summary tree of 29 fungal RALF sequences and five plant RALF sequences was generated by pruning each tree in the posterior distribution of RALF gene trees to just these 34 tips, and then producing an extended majority-rule tree from the posterior using PAUP 4.0 (<http://paup.csit.fsu.edu/>). The resulting summary tree was rooted and ladderized using TreeGraph 2 (Stöver and Müller, 2010). The full summary tree of all sequences was generated using the same method, but without pruning. To produce a combined phylogeny and alignment, the RALF sequences, including non-conserved residues, were manually aligned, re-ordered to match the pruned summary tree and individual amino acid residues coloured using JalView (Waterhouse *et al.*, 2009).

Fox RALF homologue sequences were aligned in Geneious 7.1.5 (Biomatters, Auckland, New Zealand) using MUSCLE alignment (Edgar, 2004).

Synthetic peptides

Four synthetic RALF-like peptides were produced by Mimotopes (Notting Hill, Australia) (Fig. S4). First, synthetic tomato (*Solanum lycopersicum*) RALF-like (SI-RALF) peptide (9.2 mg with minimum purity of 96%) was chosen as a positive control because of its similar domain structure to the RALF peptides described by Pearce *et al.* (2010) in a study describing important residues for activity and for its similarity to *Fol* race 2. The synthetic peptide started at the first amino acids reported to be required for RALF activity ('YISY') (Pearce *et al.*, 2010). Second, inactive SI-RALF peptide (SI-RALFΔ) was used as a negative control peptide (9.4 mg with minimum purity of 82%). This peptide was designed missing the YISY motif required for RALF function (Pearce *et al.*, 2010). We synthesized two peptides of fungal origin: *Fol* race 2 RALF (RALF-B) (9.3 mg with minimum purity of 86%) and *Fol* race 3 RALF (RALF-C) (9.2 mg with minimum purity of 77%). These two *Fox* RALF peptide sequences are representative of the two main RALF variants observed in *Fox* (Fig. 3). All peptides were synthesized in oxidized form, with disulphide bonds between cysteine residues at the indicated positions (Figs 2 and 3).

Root inhibition in response to fungal RALF synthetic peptides

Tomato seeds (*Solanum lycopersicum* cv. MoneyMaker) were surface sterilized for 10 min in 4% sodium hypochlorite–10% ethanol and germinated in 2 mL of 50% Murashige and Skoog (MS) medium on an orbital shaker for 2 days. Three lots of three germinating seeds were placed into 2 mL of fresh 50% MS medium with a final concentration of 10 μM synthetic peptide and incubated on the orbital shaker for 2 days. After this time, seedling weight was measured and comparisons in developmental morphology were made. Tukey's test was used to determine which differences between treatments were significant for all pairwise combinations of treatments, and to correct for multiple testing.

Measurement of ROS, induction of alkalinization and MAPK production in tomato and *N. benthamiana* in response to synthetic RALF peptides

ROS assays were performed as described previously (Segonzac *et al.*, 2011), except that L-012 (Wako Chemicals, Cape Charles, USA) was used instead of luminol, and luminescence was measured on a Tecan plate reader, Infinite M200 PRO (Tecan, Männedorf, Switzerland). MAPK

activation assays were performed as described previously (Segonzac *et al.*, 2011). For the alkalization experiment, tomato or *N. benthamiana* leaves were cut the day before the assay in strips of approximately 1 cm × 0.1 cm, and incubated in double-distilled H₂O (ddH₂O) overnight. The next day, 12–15 leaf strips of each were transferred to one well of a 12-well plate containing 2 mL of sterile ddH₂O. Leaves were incubated in the absence or presence of 1 μM RALF peptide for 1 h with shaking at 120 rpm. The pH was measured using a pH electrode until the reading was stable. The graphs show the average of three biological replicates and its estimated standard error. Different letters depict significant differences using Tukey–Kramer honest significant difference (HSD) test with $P < 0.01$.

RT-PCR analysis of RALF-B expression during infection

Two-week-old susceptible tomato cv. M82 seedlings were inoculated by dipping their roots in a suspension of 1×10^7 conidia/mL of *Fol* isolate #1943, or mock inoculated by dipping in water. *Fol*-inoculated and mock-inoculated plants were then grown in a controlled-environment growth room with a 25 °C, 16-h day/20 °C, 8-h night cycle until collection of the samples. Roots of three to four *Fol*-infected or mock-inoculated plants were collected at 3 and 6 dpi, washed with sterile deionized water, pooled in a microcentrifuge tube and frozen in liquid nitrogen ready for RNA extraction. Frozen root samples were ground in liquid nitrogen and total RNA was extracted using a Plant RNeasy kit (Qiagen, Hilden, Germany) following the manufacturer's instructions. Total RNA (2 μg) was treated with 2 μL of RQ1 RNase-Free DNase (Promega, Madison, WI, USA) in a reaction volume of 20 μL containing 1 × RQ1 DNase reaction buffer (400 mM Tris-HCl, pH 8.0, 100 mM MgSO₄, 10 mM CaCl₂; Promega) and 1 μL of RNasin ribonuclease inhibitor (Promega). The reaction was incubated at 37 °C for 30 min, followed by an inactivation step at 65 °C for 20 min. Treated RNA was reverse transcribed into cDNA using Superscript III Reverse Transcriptase (Invitrogen, Carlsbad, CA, USA) and an oligo [dT]_{12–18} primer (Invitrogen) following the manufacturer's instructions. PCR (35 cycles) was carried out using Phire Hot Start II DNA Polymerase (Thermo Fisher Scientific, Waltham, MA, USA) in a reaction volume of 25 μL containing 1 μL of cDNA template according to the manufacturer's instructions. Primers 5'-GCTGAAGCAACCCCTATAA-3' and 5'-TTACGATCCGGTTACCAAGC-3' were used to amplify RALF*Fol*, and primers 5'-AGCCTTACACCATCCGCTAC-3' and 5'-CGCTGTAGTTGACCTACCA-3' were used to amplify *FEM1* (*Fusarium* extracellular matrix protein 1) (Schoffelmeyer *et al.*, 2001) as a positive control for fungal gene expression. RNA and genomic DNA (50 ng) from non-inoculated tomato plants and 5-day-old cultures of *Fol* isolates #1943 and 4287 were used as controls. Controls also included reactions in which reverse transcriptase or template DNA was omitted.

Fungal strains and pathogenicity test on tomato

The tomato M82 cultivar susceptible to *Fol* race 3 was used for pathogenicity tests. Australian *Fol* race 3 isolate #1943 was used to generate RALF knockout mutants. Pathogenicity tests were performed by the root dip method (Mes *et al.*, 1999; Wellman, 1939). Using this method, roots of 3-week-old plants of the tomato cultivar M82 (susceptible to *Fol* race 3) were wounded and subsequently inoculated with *Fol* race 3. Disease symptoms were scored based on plant fresh weight and disease index, as described by Rep *et al.* (2004).

Construction of *Fol* RALF deletion vectors

Flanking regions of the RALF gene were amplified from *Fol* race 3 genomic DNA using primers RALF_FR1 and RALF_FR2 for the upstream flanking region and RALF_FR3 and RALF_FR4 for the downstream flanking region (Table S2, see Supporting Information). The upstream flanking region was cloned via *KpnI* and *BsrGI* restriction sites into the 5' end of the hygromycin resistance cassette in the binary vector pPK2HPH (Michielse *et al.*, 2009), and the downstream flanking region was cloned via *XbaI* and *HindIII* sites into the 3' end of the hygromycin resistance cassette to generate pPK2HPH:ΔRALF.

A modified *HSVtk* gene cassette was generated and used as a second selection marker, as described by Khang *et al.* (2005). Briefly, the promoter of the *trpC* gene of *Aspergillus nidulans* was amplified from the pGpdGFP binary vector (Sexton and Howlett, 2001) using the primers *trpC_p-F* and *trpC_p-R* (Table S2), and cloned into the pGEMt easy vector (Promega). The *HSVtk* gene was obtained from DNA of the herpes simplex virus (HSV) isolate UL23 (provided by Professor David Tschärke, The Australian National University, Canberra, Australia) using the primers *HSVtk-F* and *HSVtk-R* (Table S2), and cloned into the pGEMt:trpC promoter intermediate vector via *SpeI* and *SacI* sites. The terminator sequence of the *Fol* β-tubulin gene was obtained from *Fol* genomic DNA using the primers βtub-F and βtub-R, and cloned into the pGEMt:trpC promoter:HSVtk coding sequence intermediate vector via *SacI* and *PmeI* restriction sites. The entire *HSVtk* cassette was then transferred to the pPK2HPH:ΔRALF vector and placed next to the RALF downstream flanking region, using *HindIII* and *PmeI* sites (Fig. S6A).

Fol transformation was performed using the *Agrobacterium tumefaciens* strain LBA4404 containing the appropriate binary vector. The transformation protocol was adapted from Mullins *et al.* (2001) and Takken *et al.* (2004). The selection medium for the RALF knockout mutants was supplemented with 75 μg/mL augmentin, 50 μg/mL hygromycin and 5 mM 5-fluoro-2'-deoxyuridine (F2dU). Deletion of the RALF gene was confirmed by PCR using the primers *scRALF_5'-F*, *RALF_int-R*, *RALF_int-F*, *scRALF_3'-R* and *scGDP-R* (Table S2 and Fig. S6B).

NOTE ADDED IN PROOF

Consistent with our findings, Vlaadingerbroek *et al.* (2016; Mol. Plant Pathol. DOI: 10.1111/mpp.12440) have shown, using the same kind of tomato pathogenicity assay and scoring system we used, that complete loss of *Fol* chromosome 12, which carries the RALF-B gene, has no significant effect on *Fol* race 2 pathogenicity.

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SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article at the publisher's website:

Fig. S1. Gene tree of all identified rapid alkalization factor (RALF) sequences from plants and fungi. This tree was inferred using the RALF peptide domain amino acid sequences from 433 plant RALF sequences and 29 fungal RALF sequences (highlighted in bold). Support values on each branch refer to the percentage posterior probability of each clade. Fungal RALF sequences are labelled with the scientific name of the fungal species or isolate. Plant RALF sequences are labelled with their NCBI GI code, binomial name and NCBI RefSeq description. The tree was rooted to split the tree into RALF 27-like sequences and other RALF sequences, a split with 100% posterior support.

Fig. S2. There are three divergent rapid alkalization factor (RALF) groups within the Botryosphaeriaceae (A). Particular

motifs are unique among these groups (A). In particular, Group B1 has a motif shared between few other species (plants and fungi) (B). One of *Macrophomina phaseolina*'s Group B11 RALFs shows 100% identity for the mature peptide region with species of soy. These are primary hosts of *M. phaseolina*.

Fig. S3. A limited number of bacteria encode a protein with a pertussis toxin subunit (S1) and a C-terminal rapid alkalization factor (RALF) domain (A). This C-terminal RALF domain in these bacteria shares homology with plant species RALFs (plant species outlined) (B).

Fig. S4. Amino acid sequences of peptides synthesized for the reactive oxygen species (ROS) burst and mitogen-activated protein kinase (MAPK) activation assays

Fig. S5. Concentration-dependent rapid alkalization factor (RALF)-mediated production of reactive oxygen species (ROS) in *Nicotiana benthamiana* leaves. (A) ROS burst over time on treatment with 1, 10, 50, 100 and 500 nM RALF-B. (B) ROS burst over time on treatment with 100 nM, 1 μM and 10 μM of RALF-C.

Fig. S6. Activation of mitogen-activated protein kinases (MAPKs) in *Nicotiana benthamiana* leaves at 5 and 15 min after treatment with the synthetic rapid alkalization factor (RALF) peptides as indicated.

Fig. S7. Generation and screening of *Fil* race 3 transformants containing a knockout of the *RALF* gene (Δ RALF). (A) Schematic representation of the *RALF* gene replacement using the hygromycin phosphotransferase (*hph*) gene as a selectable marker and the thymidine kinase (*tk*) gene as a counter-selectable marker. Arrows indicate primers pairs used for screening. (B) Polymerase chain reaction (PCR) detection of *RALF* gene using scRALF 5'-F/RALF_int-R primers (primer set B; left gel image) and RALF_int-F/scRALF 3'-R primers (primer set C; right gel image). Both gel images indicate the absence of the *RALF* coding sequence in the four Δ RALF transformants. Fol, *Fusarium oxysporum* f. sp. *lycopersici* DNA. (C) PCR with scRALF 5'-F/sc_gdp-R primers (primer set D) indicates the presence of the *hph* cassette in the Δ RALF transformants. Ect, DNA from a transformant with an ectopic T-DNA insertion in *Fol*. (D) Pathogenicity test on M82 plants with *Fol* wild-type race 3 (WT), a *Fol* transformant with an ectopic T-DNA insertion or *Fol* Δ RALF transformants Δ 23 and Δ 24. Photographs were taken at 21 days post-inoculation (dpi). (E) Distribution of disease scores for the symptoms observed in two experiments for 10 plants for each treatment at 21 dpi. Probability values were obtained using the one-tailed, non-parametric, Mann–Whitney test to determine significant differences ($P \leq 0.05$) between treatments.

Table S1. Arabidopsis RALF sequences used as BLAST queries.

Table S2. Primers used in this study.