

FcRav2*, a gene with a ROGDI domain involved in Fusarium head blight and crown rot on durum wheat caused by *Fusarium culmorum

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SUMMARY

Fusarium culmorum is a soil-borne fungal pathogen which causes foot and root rot and Fusarium head blight on small-grain cereals, in particular wheat and barley. It causes significant yield and quality losses and results in the contamination of kernels with type B trichothecene mycotoxins. Our knowledge of the pathogenicity factors of this fungus is still limited. A transposon tagging approach based on the *mimp1/impala* double-component system has allowed us to select a mutant altered in multiple metabolic and morphological processes, trichothecene production and virulence. The flanking regions of *mimp1* were used to seek homologues in the *F. culmorum* genome, and revealed that *mimp1* had reinserted within the last exon of a gene encoding a hypothetical protein of 318 amino acids which contains a ROGDI-like leucine zipper domain, supposedly playing a protein–protein interaction or regulatory role. By functional complementation and bioinformatic analysis, we characterized the gene as the yeast Rav2 homologue, confirming the high level of divergence in multicellular fungi. Deletion of *FcRav2* or its orthologous gene in *F. graminearum* highlighted its ability to influence a number of functions, including virulence, trichothecene type B biosynthesis, resistance to azoles and resistance to osmotic and oxidative stress. Our results indicate that the FcRav2 protein (and possibly the RAVE complex as a whole) may become a suitable target for new antifungal drug development or the plant-mediated resistance response in filamentous fungi of agricultural interest.

Keywords: fungal pathogens, fungicide, *Fusarium graminearum*, Fusarium head blight, molecular target, transposon tagging, virulence genes.

INTRODUCTION

Fusarium culmorum (W.G. Smith) Sacc., together with *F. graminearum* Schwabe and *F. pseudograminearum* O'Donnell & T. Aoki, are considered to be the most devastating fungal pathogens on small-grain cereals (soft and durum wheat, barley, oat, rye and triticale). The interest in these species is justified by their role in the onset of two distinct diseases, namely foot and root rot (FRR, also known as 'crown rot') and Fusarium head blight (FHB) (Goswami and Kistler, 2004; Kazan *et al.*, 2012; Scherm *et al.*, 2013; Wagacha and Muthomi, 2007; Xu and Nicholson, 2009). FRR infection results in pre- or post-emergence seedling death, brown discoloration on coleoptiles and the formation of whiteheads, leading to significant yield losses. However, yield and grain quality are particularly affected when these pathogens induce FHB, infecting the heads at anthesis and colonizing tissues until grain harvest. FHB infection causes contamination of the grain with mycotoxins, such as type B trichothecenes, i.e. sesquiterpene epoxides that are able to inhibit eukaryotic protein synthesis and induce apoptosis, and may play an important role as virulence factors.

Significant progress has been made during recent years towards a better understanding of the processes involved in FHB, especially in *F. graminearum* (Fu *et al.*, 2013; Jiang *et al.*, 2010; Kazan *et al.*, 2012; Liu *et al.*, 2014; Lysøe *et al.*, 2011; Ma *et al.*, 2013; Sperschneider *et al.*, 2015). On the contrary, our knowledge on *F. culmorum* pathogenicity is still poorly understood and, although genes have been reported from whole-genome sequencing projects (Moolhuijzen *et al.*, 2013; Urban *et al.*, 2016), only a few new potential fungicide targets in this species have been reported so far (Baldwin *et al.*, 2010; Pasquali *et al.*, 2013; Scherm *et al.*, 2011; Skov *et al.*, 2004).

With the aim to identify new *F. culmorum* genes playing a role in both FHB and FRR on durum wheat, we performed a transposon-mediated mutagenesis approach based on the heterologous element *mimp1* (Hua-Van *et al.*, 2000). *mimp1* does not encode a transposase gene, but can be trans-mobilized by the

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transposase of the *impala* element (Dufresne *et al.*, 2007, 2008; Spanu *et al.*, 2012).

Here, we report the identification of a new *F. culmorum* gene (*FcRav2*) by *mimp1*-mediated insertional mutagenesis. The putative role of *FcRav2* was determined in *F. culmorum* and, as a comparison, in *F. graminearum* by the analysis of the effect of deletion on the fungal phenotype.

RESULTS

Molecular characterization of the *mimp1*-tagged mutant R38

Polymerase chain reaction (PCR) and Southern blot analyses confirmed that the excision event in the revertant strain R38 was followed by the reinsertion of the *mimp1/impala* construct into a different genome site (not shown). Based on the left flanking sequence of the *mimp1* element, BLASTN results revealed that *mimp1* had reinserted at position 6542003 of the UK99 genome draft (LT598662.1) in the gene FCUL_11566.1 located in the fourth chromosome of *F. culmorum* (Fig. S1, see Supporting Information).

Characterization of the *FcRav2* gene

FcRav2 (in *F. graminearum*, *FgRav2*, corresponding to Fg09428/FGSG_17209) has two introns and three exons, and codes for a hypothetical protein of 318 amino acids, with a molecular weight of 34.8 kDa and an isoelectric point of 6.43. The amino acid sequence contains a ROGDI leucine zipper domain (pfam10259), including a region of 30 amino acids with leucine repeats every seven or eight residues (Fig. S2, see Supporting Information). Homologous genes with unknown function were found in *F. oxysporum* (FOX_06114) and *F. verticillioides* (FVEG_03978) with sequence homologies of up to 83%, as well as in other filamentous fungi (Fig. S3, see Supporting Information). Orthology conservation was strongly confirmed (e-value EggNog e-145 in the Class *Leotiomyces*). Orthology classification from the National Center for Biotechnology Information (NCBI) allocated the gene within the Subclass Sordariomycetidae including the *Neurospora crassa* (NCU08091) and *Magnaporthe oryzae* (MGG_02604) ROGDI-containing group. A lower level of conservation was found within the Class Saccharomycetes using both EggNog and Inparanoid. As a result of the unique domain and homology along the overall protein length, we hypothesize that orthology also occurs with the *Rav-2* homologue in *Saccharomyces cerevisiae*, as well as with higher eukaryotes (Dawson *et al.*, 2008). Protein localization is probably nuclear according to WoLF-PSORT analysis (<http://www.genscript.com/wolf-psort.html>; Horton *et al.*, 2007).

According to TMpred (http://www.ch.embnet.org/software/TMPRED_form.html; Hoffmann and Stoffel, 1993), the protein may contain a transmembrane domain (AA 178–201) with its N-

terminus in the internal membrane region. The hypothetical structure obtained with RaptorX (Källberg *et al.*, 2012) is reported in Fig. S4 (see Supporting Information).

Generation of *FcRav2/FgRav2* deletion mutants in *F. culmorum* UK99 and *F. graminearum* PH1, and annotation confirmation

Seven hygromycin B-resistant transformants were obtained from *F. culmorum* UK99, with five showing a distinct deletion and two being ectopic transformants (Fig. 1). One ectopic strain (*FcB6*) and two $\Delta FcRav2$ strains ($\Delta FcRav2$ B24, $\Delta FcRav2$ B51) were used for all the following assays and characterizations. In addition, five FGSG_17209 deletion mutants were obtained from *F. graminearum* PH1, with $\Delta FgRav2$ G8 and $\Delta FgRav2$ G10 (Fig. 1) selected for further analyses.

To verify the *in silico* annotation of the gene, the wild-type strain UK99 was treated with 11 μ mol of bafilomycin, which is known to inhibit specifically the V-ATPase complex (Faraco *et al.*, 2014). Indeed, the phenotype of the wild-type treated with bafilomycin reproduced the same morphological features as the mutants, including colony morphology, hyphal hyperbranching and pigmentation (Fig. 2), allowing confirmation that at least part of the phenotypic effect of the *FcRav2* mutation can be mimicked by a V-ATPase-inhibiting drug.

Deletion of the *FcRav2/FgRav2* gene involves significant changes in the physiological and metabolic profiles of *F. culmorum* and *F. graminearum*

All *FcRav2* deletion mutants, as well as the revertant strain R38, showed significantly decreased growth compared with the recipient strains *FcM7*, *F. culmorum* UK99 and *F. graminearum* PH1 when coping with multiple osmotic stresses, whereas the ectopic transformant strain *FcB6* displayed the same responsiveness as the *F. culmorum* wild-type strain UK99 (Table 1; Figs S5–S8, see Supporting Information). The addition of 2 M sorbitol or 0.02% (w/v) sodium dodecylsulfate (SDS) led to more significant growth reductions than did 1 M NaCl. *Fusarium graminearum* was completely inhibited at 0.02% (w/v) SDS, and therefore PH1, $\Delta FgRav2$ G8 and $\Delta FgRav2$ G10 were characterized in a second test series on 0.01% (w/v) SDS, where deletion mutants displayed a significant growth reduction relative to the wild-type PH1 (Table 1).

Only *F. culmorum* strain *FcM7* and its revertant R38 were able to grow in the presence of 30 mM $K_2S_2O_8$, with R38 being significantly inhibited (over 70%) relative to *FcM7*. In the presence of 20 mM $K_2S_2O_8$, all *F. culmorum* and *F. graminearum* deletion mutants were significantly inhibited relative to the respective wild-type (Table 1).

Conidial germination was significantly impaired in the *F. culmorum* and *F. graminearum* deletion mutants relative to the wild-type strains. In contrast, the germination capability of the

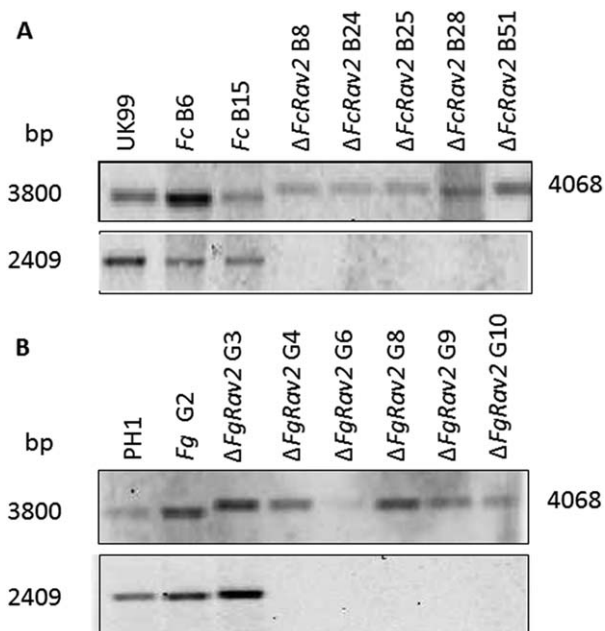


Fig. 1 Southern blot analysis of *Fusarium culmorum* (A) and *F. graminearum* (B) transformants. Top: *EcoRV*-digested DNAs were blotted and hybridized with a gene upstream probe obtained with primers 1F/2R (expected size for wild-type and ectopic transformant, 3800 bp; expected size for deletion mutants, 4068 bp). Bottom: *BglIII*-digested DNAs were blotted and hybridized with an internal *FcRav2* gene upstream probe obtained with primers NF/NR (expected size for wild-type and ectopic transformant, 2409 bp).

revertant strain R38 was not reduced relative to that of the co-transformant strain *FcM7* (Table 1).

FcRav2 deletion dramatically reduced mycelium hydrophobicity: whereas, for both the wild-type strain UK99 and the ectopic transformant *FcB6*, the time required to adsorb a 20- μ L drop of

water was more than 15 min, in $\Delta FcRav2$ B24 and $\Delta FcRav2$ B51, the water droplet was immediately adsorbed on deposition (data not shown).

Tebuconazole sensitivity is significantly increased in the *mimp1*-tagged mutant and in deletion mutants

The addition of 0.5 μ g/mL of tebuconazole to Czapek dox agar produced a significant reduction in colony growth in all deletion mutants. In *F. culmorum*, growth of the *mimp1*-tagged mutant R38 was decreased by 70% compared with the co-transformant *FcM7*, whereas *FcRav2* deletion resulted in 25%–35% growth inhibition on fungicide-amended medium. Similarly, *F. graminearum* deletion mutants showed 50% growth reduction compared with the *F. graminearum* wild-type strain PH1 in the presence of tebuconazole (Table 1).

***FcRav2* gene plays a major role in sugar metabolism**

Using a phenotype microarray approach, *F. culmorum* wild-type strain UK99, its deletion mutant $\Delta FcRav2$ B24 and the ectopic transformant strain *FcB6* were further screened for their putative phenotype differences associated with low-molecular-weight carbon uptake and metabolism. The complete set of triplicate OD₇₅₀ (optical density at 750 nm) readings recorded every 15 min from 0 to 72 h of incubation on 96 different carbon sources is reported in Table S1 (see Supporting Information).

Based on the analysis of the growth curves, the interval between 60 and 72 h was selected as the most informative to highlight differences in growth amongst the three tested strains. The heat map depicted in Fig. 3 reports the average OD₇₅₀ readings recorded during the 60–72-h interval on the 30 most differentiating carbon sources, listed in order of decreasing proportional

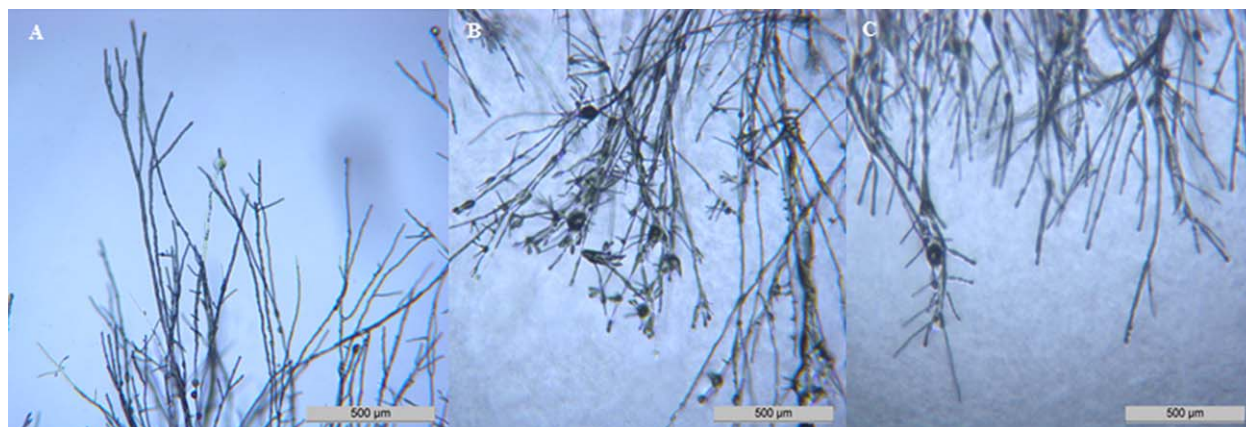


Fig. 2 Comparison of *Fusarium culmorum* $\Delta FcRav2$ mutant phenotype and *in vitro* effect of the vacuolar type H(+)-ATPase inhibitor bafilomycin A1. (A) *Fusarium culmorum* wild-type strain UK99 colony border after 48 h of growth on Czapek dox agar. (B) UK99 grown on Czapek dox agar containing 11 μ mol of bafilomycin A1. (C) *Fusarium culmorum* mutant $\Delta FcRav2$ B51 grown on Czapek dox agar. Microscopic detail (40 \times) shows hyperbranching and thickening of the hyphal tips in the $\Delta FcRav2$ mutant and in the wild-type exposed to bafilomycin A1.

Table 1 Phenotype assay of the strains used in this study.

Strain	Sorbitol (2 M)	NaCl (1 M)	SDS (0.02%)	SDS (0.01%)	K ₂ S ₂ O ₈ (30 mM)	K ₂ S ₂ O ₈ (20 mM)	Tebuconazole (0.5 µg/mL)	Germinated conidia (%)
<i>FcM7</i>	2.30 ± 0.04	5.26 ± 0.02	2.14 ± 0.02	N.D.	2.56 ± 0.14	N.D.	5.26 ± 0.02	0.74 ± 0.01
R38	0.84 ± 0.02**	0.82 ± 0.02**	1.14 ± 0.02**	N.D.	0.70 ± 0.0**	N.D.	1.58 ± 0.02**	0.77 ± 0.01
<i>Fusarium culmorum</i> UK99	2.98 ± 0.02	4.88 ± 0.05	2.78 ± 0.02	N.D.	No growth	4.87 ± 0.07	4.13 ± 0.14	0.30 ± 0.02
<i>FcB6</i> (ectopic)	2.68 ± 0.10	5.02 ± 0.05	2.90 ± 0.03	N.D.	No growth	5.10 ± 0.06	4.33 ± 0.05	0.28 ± 0.01
$\Delta FcRav2$ B24	1.50 ± 0.07**	1.40 ± 0.03	1.80 ± 0.03**	N.D.	No growth	2.50 ± 0.0**	2.77 ± 0.03**	0.05 ± 0.01**
$\Delta FcRav2$ B51	1.56 ± 0.05**	1.80 ± 0.09	1.86 ± 0.02**	N.D.	No growth	2.77 ± 0.03**	3.13 ± 0.05**	0.02 ± 0.01**
<i>Fusarium graminearum</i> PH1	4.02 ± 0.04	4.86 ± 0.04	No growth	4.83 ± 0.03	No growth	5.00 ± 0.0	5.10 ± 0.10	0.34 ± 0.02
$\Delta FgRav2$ G8	1.08 ± 0.06**	2.84 ± 0.05**	No growth	3.26 ± 0.03**	No growth	3.97 ± 0.07**	2.08 ± 0.08**	0.14 ± 0.01**
$\Delta FgRav2$ G10	1.6 ± 0.09**	3.78 ± 0.05**	No growth	3.13 ± 0.08**	No growth	3.70 ± 0.15**	2.36 ± 0.05**	0.15 ± 0.01**

Colony growth was measured after 3 days of incubation at 25 °C on Czapek dox agar supplemented with 2 M sorbitol, 1 M NaCl, 0.01%–0.02% (w/v) sodium dodecylsulfate (SDS) (osmotic stress), 20–30 mM K₂S₂O₈ (oxidative stress) or 0.5 µg/mL tebuconazole (fungicide sensitivity). Percentage spore germination was evaluated after 4 h of incubation in Czapek dox broth. Three independent tests were performed with three replicates for each test.

Results are expressed as colony diameter growth (cm) ± standard error. Significantly different from the control (***P* < 0.01) based on Dunnett's test.

N.D., not determined.

growth difference between the deletion mutant $\Delta FcRav2$ B24 and its wild-type strain UK99.

The deletion mutant $\Delta FcRav2$ B24 was impaired in its ability to catabolize different mono-, di- and trisaccharide carbon sources (particularly L-sorbose, D-xylose, α -methyl-D-glucoside, lactulose, L-arabinose, L-fucose, palatinose, D-ribose, D-mannose, D-fructose, D-fucose, D-raffinose, D-melezitose), with percentage growth rates between 45% and 70% relative to the wild-type UK99 and the ectopic strain *FcB6* (Fig. 3). Moreover, the deletion mutant grew at a reduced rate on 2-amino ethanol and on weak acids, such as bromosuccinic acid and β -hydroxy-butyric acid (Fig. 3).

***FcRav2* influences the severity of crown rot and FHB on durum wheat**

In a preliminary screening, the *mimp1*-tagged mutant R38 was compared with the co-transformant strain *FcM7* and the wild-type strain UK99 to evaluate the effect on seed germination. Abundant mycelium formation by all tested strains was observed after 24–48 h. However, only mutant R38 did not hamper the emergence of the primary root from the caryopsis, whereas *FcM7* and the highly virulent strain UK99 killed 100% of the inoculated durum wheat seeds before germination (data not shown).

Glasshouse experiments further confirmed the role of *FcRav2*/*FgRav2* in both crown rot and FHB. Insertion of the *mimp1* transposable element or deletion of the *FcRav2*/*FgRav2* gene in *F. culmorum* and *F. graminearum* caused a highly significant (*P* < 0.01) reduction in crown rot symptoms on durum wheat seedlings for all tested mutants, whereas the ectopic transformant *FcB6* behaved similarly to the wild-type reference strains UK99 and PH1, which caused disease incidences of 95 to 100, respectively (Table 2; Fig. S9, see Supporting Information).

In spray inoculation tests, *FcRav2* inactivation by *mimp1* insertion (R38) or deletion ($\Delta FcRav2$ B51 and $\Delta FcRav2$ B24) reduced FHB symptoms on durum wheat heads at 21 days post-inoculation (dpi) to 10% and 5%–20% compared with *F. culmorum* control strains *FcM7* and UK99, respectively (Table 2; Fig. S9). In contrast, the ectopic transformant *FcB6* was not significantly affected in its virulence towards the host plant.

In *F. graminearum*, *FgRav2* deletion mutants completely lost their pathogenicity, hence differing significantly from the wild-type reference strain PH1 (Table 2).

Infection experiments showed a minimal, and insignificant, modulation of gene expression during the different stages of interaction with the plant. Indeed, expression profiles obtained in *F. graminearum* from array studies suggest that the gene modifies its expression according to the phenological stage of infection, but that these changes are dependent on various conditions and therefore show a high variability in experimental repetition *in planta* (Fig. 4).

***FcRav2* is involved in type B trichothecene production**

In *in vitro* experiments, only the *F. culmorum* *mimp1*-tagged mutant R38 lost completely its ability to produce type B trichothecenes, whereas trichothecene production by the deletion mutants $\Delta FcRav2$ B24 and $\Delta FcRav2$ B51 did not differ significantly from that of the wild-type strain UK99 or the ectopic transformant *FcB6* (Table 2). In the case of *F. graminearum*, mutant $\Delta FgRav2$ G8 was not significantly affected by gene deletion in its ability to produce deoxynivalenol (DON) and its acetylated form (3-ADON), whereas trichothecene production by the mutant $\Delta FgRav2$ G10 was reduced by approximately 70% (Table 2).

Carbon source	$\Delta FcRav2$ B24	Fc B6	UK99
L-sorbose	106	234	238
D-Xylose	125	267	252
2-Amino Ethanol	62	107	113
Bromosuccinic Acid	77	133	138
α -Methyl-D-Glucoside	105	186	183
Lactulose	86	147	146
L-Arabinose	118	195	194
L-Fucose	78	135	127
Palatinose	170	274	272
β -Hydroxy-butyric Acid	71	107	112
D-Ribose	133	199	198
D-Mannose	159	231	235
D-Fructose	157	241	230
D-Raffinose	176	259	253
D-Melezitose	186	253	266
m-Inositol	132	191	185
α -D-Glucose	190	265	263
D-Galactose	145	181	198
Maltose	166	226	226
Maltotriose	208	277	276
Stachyose	121	157	156
D-Melibiose	180	240	232
Adenosine	201	226	258
Sucrose	214	272	273
Gentiobiose	185	235	236
β -Methyl-D-Glucoside	169	220	211
β -Methyl-D-Galactoside	181	214	224
D-Cellobiose	228	292	277
Turanose	234	288	282
D-Mannitol	227	258	263

Fig. 3 Phenetic heat map of carbon utilization patterns of *Fusarium culmorum* deletion mutant $\Delta FcRav2$ B24 relative to its wild-type strain UK99 and to ectopic transformant strain FcB6. Average OD₇₅₀ (optical density at 750 nm) readings recorded during the 60–72-h interval on the 30 most differentiating carbon sources are listed in order of decreasing proportional growth difference between $\Delta FcRav2$ B24 and UK99.

In accordance with the development of FRR symptoms, type B trichothecene levels reached 1000–2000 ng/g in seedling stem tissue infected by *F. culmorum* strains FcM7, UK99 and FcB6 and by *F. graminearum* strain PH1. In contrast, trichothecene mycotoxins were not detected in wheat seedlings on inoculation with the *mimp1*-tagged mutant R38 or with deletion mutants of both *F. culmorum* UK99 and *F. graminearum* PH1 (Table 2).

When *F. culmorum* FcM7 was spray inoculated, multiple infection sites in the spikes resulted in high levels of trichothecene content in infected kernels, whereas the *mimp1*-tagged mutant R38 produced approximately 15% mycotoxin compared with the co-transformant strain FcM7 (Table 2). Infected kernels collected

from spikes that had been spray inoculated with *F. culmorum* ectopic transformant FcB6 contained unusually high trichothecene content (approximately 12 000 ng/g tissue), whereas the wild-type strain UK99 and both deletion mutants ($\Delta FcRav2$ B24 and $\Delta FcRav2$ B51) did not differ significantly in their DON production on kernel colonization (approximately 1700–2600 ng/g tissue; Table 2).

DISCUSSION

Within the course of an extensive transposon-mediated mutagenesis programme in the wheat pathogen *F. culmorum* (Spanu *et al.*, 2012), a pathogenicity mutant was identified among over 2000 *niaD* revertants on minimal medium containing nitrate as the sole nitrogen source. In this mutant, the *mimp1* transposable element was integrated within the last exon of the *FcRav2* gene, encoding a hypothetical protein of 318 amino acids, and containing a ROGD1-like leucine zipper domain known to have a regulatory role (pfam 10259). The ROGD1 domain is conserved in eukaryotic genomes from *Drosophila* to yeasts and is mostly unique. Our protein presents a low level of similarity with the *RAV2* gene from yeast (Seol *et al.*, 2001), part of the RAVE complex involved in the assembly and disassembly of the V-ATPase complex in yeast (Kane, 2006).

Based on the analysis of phenotype, expression and function, we hypothesize that, in *F. graminearum* and *F. culmorum*, the gene is also a *rav2* homologue. As *RAV2* gene homologues have not been investigated previously in other fungi, apart from *Schizosaccharomyces pombe*, where the gene is important for sugar response and calcium stress (Dawson *et al.*, 2008), our work represents the first characterization of the role of *RAV2* in filamentous fungi. More importantly, we showed that the protein affects the fitness of the fungus, including the ability to grow on different sugars, as well as the sensitivity to pH and fungicides, such as tebuconazole.

Treatment with the specific V-ATPase inhibitor bafilomycin reproduced the same morphological features as in $\Delta FcRav2$ mutants, including the hyphal hyperbranching observed on deletion of the *vmaA1* gene, which encodes one of the subunits of the V-ATPase in *Aspergillus nidulans* (Melin *et al.*, 2004). The V-ATPase has been shown to play a role in virulence in different fungi (Chen *et al.*, 2013; Hilty *et al.*, 2008; Patenaude *et al.*, 2013). The impairment of pathogenicity observed in *F. graminearum* and *F. culmorum* confirms indirectly the putative function of *FcRav2/FgRav2* as the homologue of *RAV2*, a controller of vacuolar and endosomal processing.

Further investigations are needed to experimentally confirm the role of the gene in filamentous fungi. Provided that the *RAV2* gene may be present only in fungi (Parra *et al.*, 2014), it appears now pivotal to focus on molecules able to interfere with *RAV2* *in planta* in order to impair the toxigenic process, or to foster the

Table 2 Production of type B trichothecenes *in vitro* and *in planta* and virulence on durum wheat (cv. Iride) of the strains used in this study.

Strain	DON production <i>in vitro</i> (ng/mL medium) [†]	Disease incidence (FRR) (0–100)	DON production in wheat seedlings (ng/g tissue) [†]	Disease incidence (FHB) (0–100)	DON production in wheat kernels (ng/g tissue) [‡]
<i>Fusarium culmorum</i> FcM7	3198.1 ± 155.5	61.9 ± 0	1092.9 ± 597.2	48.1 ± 15.1	5340.3 ± 4828.2
R38	0**	6.3 ± 5.5**	0**	4.7 ± 1.7**	900.0 ± 435.9*
<i>Fusarium culmorum</i> UK99	21 085.6 ± 2727.4	95.0 ± 6.6	2219.4 ± 1324.1	57.8 ± 7.7	2598.0 ± 0
FcB6 (ectopic)	21 612.0 ± 1060.2	96.7 ± 3.8	1826.8 ± 289.3	77.8 ± 6.9	11 979.7 ± 1952.5**
Δ <i>FcRav2</i> B24	24 044.4 ± 7431.9	22.5 ± 13.2**	0**	28.9 ± 7.0**	1732.0 ± 433.0
Δ <i>FcRav2</i> B51	22 369.5 ± 642.7	4.2 ± 5.2**	0**	22.2 ± 8.4**	2598.0 ± 1887.4
<i>Fusarium graminearum</i> PH1	6336.2 ± 831.3	100 ± 0	1316.6 ± 447.9	44.4 ± 15.0	N.D.
Δ <i>FgRav2</i> G8	7439.6 ± 743.6	6.7 ± 5.8**	0**	0**	N.D.
Δ <i>FgRav2</i> G10	2254.2 ± 349.8**	4.2 ± 5.2**	0**	0**	N.D.

Values are expressed as the mean of three or four replicates ± standard deviation.

Significantly different from the control strain (* $P < 0.05$, ** $P < 0.01$) based on Dunnett's test.

FHB, Fusarium head blight; FRR, foot and root rot; N.D., not determined.

[†]Total concentrations of deoxynivalenol (DON) and its acetylated form (3-ADON) were determined by liquid chromatography-mass spectrometry (LC-MS).

[‡]Total trichothecenes were measured by Rapid One Step Assay (ELISA lateral flow), designed for cereal flours.

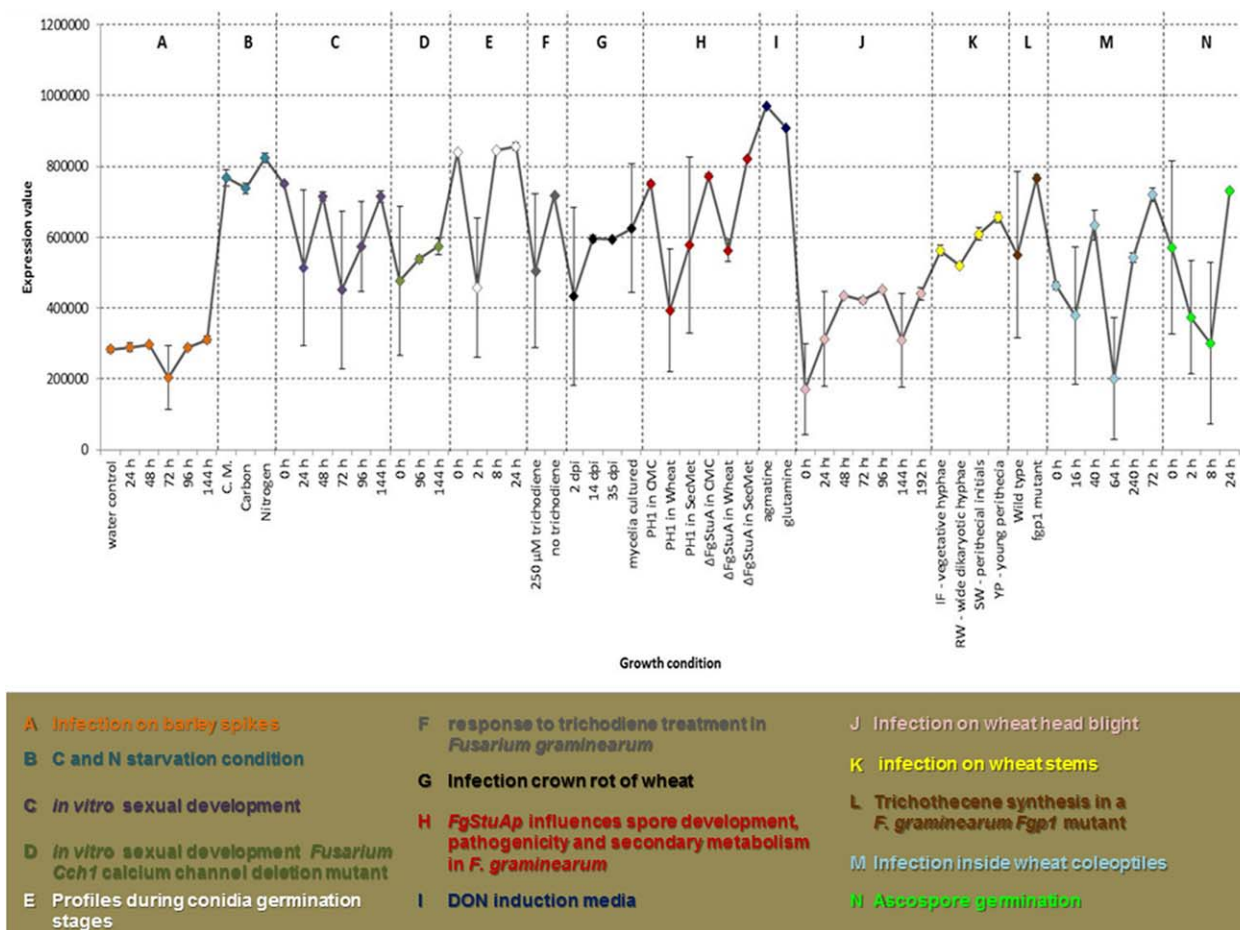


Fig. 4 Expression patterns of different Affymetrix array experiments carried out on *Fusarium graminearum* in different environmental conditions obtained from the plexdb.org platform. The results are the average of three repetitions, with bars indicating standard error. CMC, carboxy-methyl cellulose liquid medium; DON, deoxynivalenol.

activity of known fungicides, as suggested previously for human fungal pathogens (Hayek *et al.*, 2014). For this reason, we propose that RAV2 in *F. graminearum* and *F. culmorum* may represent a new target for the development of integrative approaches against fungicide adaptation and resistance, or low-sensitivity phenomena that are becoming important against different fungicides in Fusaria (Becher *et al.*, 2011; Chen and Zhou, 2009; Dubos *et al.*, 2011, 2013; Serfling and Ordon, 2014; Talas and McDonald, 2015).

FcRav2 appears to be deeply involved in a number of morphological and physiological traits, but no definite phenotypes were identified in *Candida albicans* in a large genomic screening of mutants (Noble *et al.*, 2010). Spore germination and hydrophobicity are significantly altered in deletion mutants, as is resistance to various stresses: $\Delta FcRav2$ inactivation results in increased sensitivity to osmotic and oxidative stress, and to the sterol biosynthesis inhibitor tebuconazole. The higher susceptibility to tebuconazole observed for the RAV2 mutant is a further confirmation of the importance of alternative mechanisms related to the extrusion of the compound in azole resistance. Zhang *et al.* (2010) have already hypothesized that the azole resistance mechanism also involves acidification of the medium.

Gene transcription seems to be influenced by azole treatment. Indeed, by analysis of the gene expression patterns of *F. graminearum* after treatment with sublethal concentrations of tebuconazole, Becher *et al.* (2011) found a decreased expression of FGSG_17209. This leads to the hypothesis that FGSG_17209 (*FgRav2*) may play an indirect role in controlling the expression of genes involved in compensation for azole stress, or that the gene is under-expressed as growth is decreased.

A putative role of *FcRav2* in stress resistance may also be inferred by the reduced growth of the deletion mutant on some of the tested carbon sources. For instance, L-sorbose is a well-known inhibitor of cell wall biosynthesis in fungi and shows a paramorphogenic action on *N. crassa* (Mishra and Tatum, 1972; Trinci and Collinge, 1973). Ethanolamine (2-amino ethanol) is commonly used as a wood preservative (Humar *et al.*, 2003). In addition, weak acids, such as bromosuccinic acid and β -hydroxy-butyric acid, may represent sources of stress on an impaired deleted mutant. These data suggest that *FcRav2*, as RAV2 in yeast, is involved in stress, including the stress to which *F. culmorum* is exposed when confronted with its victim plant.

The lower capacity of the *FcRav2* deletion mutant to metabolize different mono-, di- and trisaccharides in comparison with the wild-type is also concordant with the hypothesis that it represents the RAV2 homologue. Indeed, the RAVE complex in yeast is important for the reassembly of the V-ATPase machinery on sugar starvation (Smardon *et al.*, 2015). Our BIOLOG experiment confirmed the major role played by *FcRav2* in sugar metabolism. This hypothesis is corroborated by expression data obtained from independent

experiments, highlighting the under-expression of *FgRav2* in *F. graminearum* during glucose starvation (Fig. 4). In particular, among the sugars that determine the most significant differences in growth between the mutant and the wild-type, xylose and arabinose belong to the pentose and glucuronate interconversion pathway (Fungipath), which is important in conditions of glucose starvation. It is noteworthy that RAV2 is overexpressed in *S. cerevisiae* exposed to high sugar stress (Erasmus *et al.*, 2003), and that RAV2-overexpressing *Saccharomyces pastorianus* bottom-fermenting strains exhibit increased ethanol tolerance and increased fermentation rates in high-sugar medium (Hasegawa *et al.*, 2012).

As mentioned previously, the phenotype microarray analysis highlighted a significantly reduced ability of the *FcRav2* deletion mutant to utilize 2-amino ethanol as a carbon source. Interestingly, a behavioural screening of P-transposon-generated memory mutants coupled to DNA microarray analysis highlighted a *Drosophila melanogaster* ROGDI mutant and suggested its role in long-term olfactory memory (Dubnau *et al.*, 2003) and reduced ethanol tolerance (Berger *et al.*, 2008). This supports indirectly the hypothesis that homologous functions are maintained by the gene with the ROGDI domain in higher eukaryotes, although further evidence is needed to corroborate such functional homology.

FcRav2/FgRav2 have a major impact on fungal virulence in *F. culmorum* and *F. graminearum*, respectively, and influence the severity of crown rot and FHB on durum wheat. This role was clearly demonstrated by different inoculation methods (contact between mycelium and caryopsis, spray inoculation of conidia), and was confirmed in *F. graminearum*.

Infection experiments with *F. graminearum* (Güldener *et al.*, 2006; Lysøe *et al.*, 2011) showed a minimal, and insignificant, modulation of gene expression during the different stages of interaction with the plant. Indeed, expression profiles suggest that the gene modifies its expression according to the phenological stage of infection, but that these changes are dependent on different conditions probably related to the variability in experimental repetitions *in planta*.

Less convincing was the implication of *FcRav2* in mycotoxin production. The effect of *FcRav2* disruption is probably dependent on the substrate: although the *F. culmorum* *mimp1*-tagged mutant R38 lost completely its ability to produce type B trichothecenes *in vitro*, deletion mutants did not differ significantly from their wild-type strains. In seedling stem tissue, mycotoxin biosynthesis was completely inhibited on *FcRav2* deletion, whereas DON and 3-ADON were still abundantly produced by deleted strains colonizing spray-inoculated wheat spikes. This suggests that specific inducers of DON by-passing *FcRav2* regulation exist in the spike that are not able to activate the production of toxin in the stem or *in vitro*. Therefore, a plausible hypothesis is that some molecules in the spike may favour the synthesis of the toxin independent from *FcRav2* control.

Interestingly, we observed a diverse DON production *in vitro* and *in planta* for some of the strains. This phenomenon has also been observed for the transcription factor in *FgAtf1* (Nguyen *et al.*, 2013), which is also a stress regulator (Lawrence *et al.*, 2007), with opposite behaviour relative to *FcRav2*. To better elucidate the mechanisms that lead to high toxin contamination, there is a need to further characterize the specific molecules encountered by the fungus *in planta*.

In yeast, the lack of vacuolar acidification in the RAVE mutants is possibly compensated by other mechanisms of acidification of other compartments (Sardon *et al.*, 2014). This would explain the relatively mild phenotype of the *rav1Δ* and *rav2Δ* mutants. We would anticipate that this acidification process is probably under the control of *FgAtf1* in the HOG pathway. It is therefore plausible that the HOG pathway and the RAVE complex play two different, but complementary, roles in toxin production, linking toxin production and stress response pathways (Ponts, 2015). To further verify this hypothesis, we are currently investigating the cellular components of the assembly.

Expression profiles obtained from different array studies carried out on the *F. graminearum* PH1 strain confirm indirectly the homology of *FcRav2* with the *RAV-2* gene from *S. cerevisiae*. Indeed, major oscillations of the gene (higher expression) are observed in agmatine medium, which is a strong toxin inducer (Gardiner *et al.*, 2009a, b; Pasquali *et al.*, 2016b). Given the need for vacuole assembly in conditions of high toxin production (toxosomes; Menke *et al.*, 2013), it is plausible that vacuole acidification is strongly induced when high levels of toxins are produced.

From a more practical perspective, *FcRav2* is proposed to be a suitable target for new antifungal drug development or in the plant-mediated resistance response, given its pivotal role in affecting hydrophobicity, sugar metabolism, secondary metabolism, virulence and resistance to various stresses. Our work also confirms that untargeted methods for the identification of genes involved in fitness and pathogenicity are useful for genes that are not directly identified as homologous by normal orthology search programs. The further exploration of the *mimp1* reinsertion mutant library in the future will possibly allow the identification of new genes involved in the fitness of the pathogen.

EXPERIMENTAL PROCEDURES

Fungal strains and storage conditions

A *mimp1*-mediated revertant (see below) strain collection was generated from the *F. culmorum* *FcM7* co-transformant strain, integrating a single copy of the *niaD::mimp1* construct and the *impalaE* transposase gene under the constitutive control of the *gpdA* promoter (Dufresne *et al.*, 2007; Spanu *et al.*, 2012).

Homologous recombination experiments were carried out with *F. culmorum* wild-type strain UK99 (Baldwin *et al.*, 2010; kindly provided by Dr Kim Hammond-Kosack, Rothamsted Research, Harpenden, Hertfordshire,

UK) and with *F. graminearum* strain NRRL 31084 (syn. PH1; obtained from H. C. Kistler and deposited in the Luxembourg Microbial Culture Collection; Pasquali *et al.*, 2016a).

All fungal strains were routinely cultured on potato dextrose agar (PDA; Sigma-Aldrich, St. Louis, MO, USA). For long-term storage, plugs colonized by mycelium were transferred to 50% (v/v) glycerol and stored at -80°C .

Isolation of revertants obtained by the *mimp1/impala* double-component system

The *mimp1/impala* revertant strains were selected on minimal medium agar supplemented with 50 $\mu\text{g}/\text{mL}$ hygromycin (MMH50) containing sodium nitrate as sole nitrogen source (Spanu *et al.*, 2012). Briefly, selection was performed with a phenotypic assay by inoculation of 10 μL of a spore suspension [10^6 colony-forming units (CFU)/mL] of the co-transformant strain *FcM7* on MMH50 plates and by incubation at 25°C for up to 1 month. First, *nia*⁺ colonies (referred to as 'revertants') appeared 15 days after inoculation on reacquisition of nitrate reductase function, and consisted of patches of aerial mycelium with a wild-type phenotype. A single-spore culture was obtained from each collected revertant strain.

Bioassay screening of the revertant strain collection

Single-spore revertant strains were preliminarily screened for virulence during the first steps of kernel colonization using an *in vitro* bioassay described by Pasquali *et al.* (2013). Ten PDA-mycelium plugs of each revertant bearing one durum wheat seed (cv. Simeto) were placed into a 90-mm-diameter sterile Petri dish and incubated at 25°C for 3–5 days in the dark. Control assays included *FcM7*, *F. culmorum* strain UK99 and sterile PDA plugs. Inhibition of seed germination and kernel death were visually observed.

Molecular characterization of selected revertants

Excision of the *mimp1* transposable element from the *niaD* gene was evaluated by Southern blot using a *niaD*-specific probe, whereas a 120-bp fragment of the *mimp1* transposable element was used as probe to verify the reinsertion events (Table 3). Identification of the reinsertion site of *mimp1* was achieved by Splinkerette-PCR (Potter and Luo, 2010), according to the protocol described by Spanu *et al.* (2012). Flanking sequences were BLASTed to the *F. culmorum* genome (Urban *et al.*, 2016) in order to identify *mimp1* locations.

Creation of deletion mutants and mutant screening

FcRav2 deletion mutants in *F. culmorum* strain UK99 and deletion mutants for the FGSG_17209 gene homologue in *F. graminearum* strain PH-1 (*FgRav2*) were obtained by split-marker recombination, as described by Breakspear *et al.* (2011). First, screening for deletion mutants and ectopic strains was carried out by PCR with the application of several primer combinations (Table 3; 1F-FGSG17209/4R-FGSG17209, NF-FGSG17209/NR-FGSG17209, NF-FGSG17209/4R-FGSG17209 and 1F-FGSG17209/NR-FGSG17209). The PCR mix consisted of 0.5 μM of each primer, 1 \times Phusion[®] High-Fidelity PCR Master Mix with HF Buffer (New England

Table 3 Primer sequences used in this study.

Split-marker recombination	Primer sequence
1F-FGSG17209	5'-AGTCCGCTTAAGACCCTCGAC-3'
2R-FGSG17209	5'-TTGACCTCCACTAGCTCCAGCCAAGCCTAGTGAGGTGATGTTGAGATATGGA-3'
3F-FGSG17209	5'-GAATAGAGTAGATGCCGACCGGGTCCGTTGATGGAAGCGACTATTATT-3'
4R-FGSG17209	5'-GTTACACCAGTCTCGATACAACAT-3'
Identification of mutants by PCR	
1F-FGSG17209	5'-AGTCCGCTTAAGACCCTCGAC-3'
4R-FGSG17209	5'-GTTACACCAGTCTCGATACAACAT-3'
NF-FGSG17209	5'-ACTTACTAGCCATAACCTTGCCA-3'
NR-FGSG17209	5'-AGACTTCCTGAACATCATTGCTATC-3'
ITS1	5'-TCCGTAGGTGAACCTGCGG-3'
ITS4	5'-TCCTCCGCTTATTGATATGC-3'
Probes	
1F-FGSG17209	5'-AGTCCGCTTAAGACCCTCGAC-3'
2R-FGSG17209probe	5'-AAGCCTAGTGAGGTGATGTTGAGATATGGA-3'
NF-FGSG17209	5'-ACTTACTAGCCATAACCTTGCCA-3'
NR-FGSG17209	5'-AGACTTCCTGAACATCATTGCTATC-3'
Reinsertion events	
mi1	5'-TACAGTGGGATGCAATAAGTTGAATAC-3'
SacF	5'-GGCTAGATCGAGCTCCCTCCTCAG-3'
niaD144	5'-GTTTCATGCCGTGGTCGCGC-3'
niaD754r	5'-AGTTGGAATGTCCTCGTCG-3'

Biolabs, Ipswich, MA, USA) and 20 ng of DNA in a final volume of 20 μ L. A subset of putative mutants was then analysed by Southern blot with two different probes: (i) gene upstream probe obtained with primers 1F/2R (916 bp); and (ii) internal gene probe NF/NR (456 bp) (Table 3). Probe labelling, hybridization and detection reactions were carried out using the Dig High Prime DNA labelling kit and detection starter II[®] (Roche Applied Science, Basel, Switzerland), following the manufacturer's protocol.

Phenotypic analysis of transposon-tagged and deletion mutants of *F. culmorum*

The following strains were screened for altered growth under osmotic and oxidative stress or sensitivity to the fungicide tebuconazole: *mimp1*-tagged *F. culmorum* revertant strain R38, co-transformant strain FcM7, wild-type *F. culmorum* strain UK99, deletion mutants Δ Fcrav2 B24 and Δ Fcrav2 B51, ectopic transformant strain FcB6, *F. graminearum* strain PH1, and deletion mutants Δ Fgrav2 G8 and Δ Fgrav2 G10. Assays were performed on Czapek dox agar (Oxoid Limited, Basingstoke, Hampshire, UK) and Czapek dox agar supplemented with 2 M sorbitol, 1 M NaCl, 0.01–0.02% (w/v) SDS (osmotic stress), 20–30 mM K₂S₂O₈ (oxidative stress) or 0.5 μ g/mL tebuconazole (fungicide sensitivity). Ten microlitres of a titrated conidial suspension (1×10^6 CFU/mL) were spotted on the centre of each plate ($\varnothing = 60$ mm). Colony diameter growth was measured after 3 days of incubation at 25 °C in the dark, and compared with the respective controls. Conidiogenesis and spore germination were evaluated by inoculating 150 mL of Czapek dox broth with 5 mL of conidial suspension (1×10^6 CFU/mL). At 0, 2, 4 and 8 h post-inoculation (hpi) (25 °C, 150 rpm), 100 conidia were examined with a haematocytometer using a light microscope (Olympus BX41, Hamburg, Germany). Three independent tests were performed for each assay, with three replicate plates for each test.

Mycelium hydrophobicity was evaluated according to Pasquali *et al.* (2013). Briefly, a 20- μ L droplet of sterile H₂O was pipetted onto the surface of colonies grown on solid Vogel's medium (Vogel, 1956) for 5 days. The time (in seconds) needed to obtain complete droplet absorption by deletion mutants Δ Fcrav2 B24 and Δ Fcrav2 B51, and by the ectopic transformant strain FcB6, was compared with that of the wild-type strain UK99.

The wild-type *F. culmorum* strain UK99 and its deletion mutant Δ Fcrav2 B24 were further screened for phenetic differences by the application of BIOLOG FF MicroPlates (Biolog Inc., Hayward, CA, USA). The plate panel composition, which contains 95 low-molecular-weight carbon sources, is available at the BIOLOG web site (http://www.biolog.com/products-static/microbial_identification_literature.php).

Spore suspensions were prepared in carboxy-methyl cellulose liquid medium (CMC, Pasquali *et al.*, 2013) and, after two subsequent washing steps with sterile distilled water, spores were suspended in 'Inoculation fluid FF' (Biolog Inc.) at a final concentration of 1×10^4 CFU/mL. One-hundred microlitres of a spore suspension were pipetted into each of the 96 wells. The plates were incubated at 25 °C for 90 h and OD₇₅₀ was recorded every 15 min using a Microarray Omnilog Reader (Biolog, Inc.). Samples were tested in triplicate.

Pathogenicity assay on durum wheat

The *F. culmorum* co-transformant FcM7 and its revertant strain R38, wild-type strain UK99, deletion mutants Δ Fcrav2 B24 and Δ Fcrav2 B51, the ectopic transformant strain FcB6, *F. graminearum* strain PH1, and deletion mutants Δ Fgrav2 G8 and Δ Fgrav2 G10 were tested for virulence *in planta*. Both FRR and FHB were evaluated.

Mycelium plugs, each bearing one durum wheat seed (*Triticum durum* cv. Iride, kindly provided by Unità di Ricerca per la Valorizzazione Qualitativa dei Cereali, CRA-QCE, Rome, Italy), were placed in a plastic sowing

pot and covered with sterile soil. Durum wheat seeds placed onto sterile PDA plugs (1 seed/plug) served as negative controls. FRR severity was assessed after 21 days of incubation at 25 °C in a glasshouse, and evaluated using the McKinney index (McKinney, 1923; Spanu *et al.*, 2012). Three independent tests were carried out, each consisting of three replicates of 10 seedlings for each fungal strain.

The ability to cause FHB was tested on durum wheat (cv. Iride) on mid-anthesis stage wheat heads. Spray inoculation was carried out with 4 mL of each spore suspension (1×10^5 spores/mL), covering the head until runoff. Inoculated heads were covered with a transparent plastic bag for 48 h to maintain high humidity conditions. Disease incidence was evaluated every 7 days and a final score was obtained at 21 dpi using the McKinney index. Each strain was tested in four replicates of five durum wheat heads each.

***In vitro* and *in planta* mycotoxin production**

In vitro production of DON and its acetylated forms was determined in Vogel's medium (Vogel, 1956) as described by Pani *et al.* (2014), and expressed as ng/mL of culture filtrate.

The presence of trichothecenes in durum wheat seedlings (cv. Iride; three replicates of 10 seedlings each) was determined by the inoculation of seedlings at emergence with a conidial suspension (100 µL of 1×10^6 CFU/mL) of each strain. After 15 dpi, the basal portions of 10 seedlings were pooled, dried at 80 °C for 24 h, finely ground in a mortar and weighed. Samples were purified by MycoSep[®] 227 Trich⁺ columns (Romer Labs, Tulln, Austria), as described by the manufacturer, prior to liquid chromatography-mass spectrometry (LC-MS) analysis. Quantitative determinations were carried out as described previously (Pani *et al.*, 2014), using a model HP 1100 LC-MS detector (Agilent Technologies, Palo Alto, CA, USA).

The content of trichothecenes in durum wheat seeds (cv. Iride) harvested from the spray-inoculated heads was determined using the ROSA[®] FAST5 DON assay (Rapid One Step Assay, Charm Sciences, Inc., Lawrence, MA, USA), according to the manufacturer's protocol. Total trichothecene content is expressed in ng/g plant tissue.

Growth on H(+)-ATPase inhibitor bafilomycin

Fusarium culmorum wild-type strain UK99 and its mutant $\Delta FcRav2$ B51 were grown at 25 °C on Czapek dox agar amended with 6.7 µg/mL (equivalent to 11 µmol) of the vacuolar-type H(+)-ATPase inhibitor bafilomycin and on Czapek dox agar, respectively. After 48 h, colony diameters were measured and the phenotypes of the treated wild-type and untreated mutant were compared.

Bioinformatic and statistical analysis

A one-way analysis of variance, followed by multiple comparison using Dunnett's test, was performed on all data obtained from phenotypic, pathogenicity and trichothecene production assays using the Minitab[®] for Windows release 12.1 software.

Phenetic patterns were acquired with OmniLog-OL_PM_FM/Kin 1.30 and OmniLog-OL_PM_Par 1.30 software. Data were analysed separately by well with a mixed linear model that included the fixed effects of strain, sampling time, their interaction and the random effect of the replicates.

The model was solved using PROC MIXED of SAS software (SAS Institute, 2008).

Sequences for alignment were obtained from FungiDB (<http://FungiDB.org>). Sequence analyses and expression profiles were analysed using CLC Main Workbench v 7.0. Expression profiles were obtained from the PLEXdb database (www.plexdb.org; Dash *et al.*, 2012). Orthology was calculated with EggNOG 4.5 (Jensen *et al.*, 2008) and Inparanoid 8 (Sonnhammer and Östlund, 2015). The putative protein structure was generated with RaptorX structure (raptorx.uchicago.edu).

Note by the authors

Whilst this article was under revision, a paper by Nguyen *et al.* (2017) showed the effect of deletion of the homologous FcRav2 in *N. crassa*, consisting of a decrease in growth of hyphae. The gene was selected as it has been proposed to be part of a set of candidate genes with a role in multi-cellular complexity, being conserved in complex eukaryotes. Our orthology search confirmed the work by Nguyen *et al.* (2017), suggesting that FcRav2 homologues in Pezizomycotina may have many roles in governing cellular complexity that are partially lost in yeast. Moreover, the gene localization in *N. crassa*, shown by Nguyen *et al.* (2017), is consistent with our bioinformatic predictions, which suggest a cytoplasmic localization with a transmembrane domain.

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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 *FcRav2* gene structure with sequence from ENSEMBLE modified to show the insertion site of *mimp1* in the third exon of the gene. Untranslated region (UTR), introns and exons are reported.

Fig. S2 Protein domains identified in FcRav2 and the typical ROGDI leucine zipper pfam domain aligned to the FcRav2 amino acid sequence.

Fig. S3 Muscle alignment of FcRAV2 homologues.

Fig. S4 RaptorX putative three-dimensional structure of the FcRAV2 protein and summarized results of prediction probabilities for modelling. Probably, only one part of the protein is correctly modelled, given the low GDT (Global Distance Test) value (<50).

Fig. S5 Phenotype of *Fusarium culmorum mimp1* revertant R38, nit⁻ recipient and M7 co-transformant on different stress-inducing media. SDS, sodium dodecylsulfate.

Fig. S6 Phenotype of *Fusarium culmorum mimp1* revertant R38, nit⁻ recipient and M7 co-transformant on Czapek's medium amended with K₂S₂O₈ (30 mM).

Fig. S7 Phenotype of *Fusarium culmorum* UK99, FcB6 (ectopic transformant), $\Delta FcRav2$ B24 and $\Delta FcRav2$ B51 on different stress-inducing media after 10 days of growth at 25 °C.

Fig. S8 Phenotype of *Fusarium graminearum* PH1, $\Delta FgRav2$ G8 and $\Delta FgRav2$ G10 on different stress-inducing media after 10 days of growth at 25 °C. SDS, sodium dodecylsulfate.

Fig. S9 Left: foot and root rot (FRR) symptoms on 21-day-old durum wheat cv. Iride seedlings mock inoculated (A) or infected with *Fusarium culmorum mimp1* revertant R38 (B), co-transformant M7 (C) or nit⁻ recipient strain (D). Right: spikes of durum wheat cv. Iride mock inoculated (A) or infected with *mimp1* revertant R38 (B), co-transformant M7 (C) or nit⁻ recipient strain (D) at 21 days post-inoculation.

Table S1 Phenotypic growth raw data [OD₇₅₀ (optical density at 750 nm) readings] of *Fusarium culmorum* wild-type strain UK99, its deletion mutant $\Delta FcRav2$ B24 and the ectopic transformant strain FcB6 recorded in triplicate every 15 min from 0 to 72 h of incubation on 96 different carbon sources. The complete set of data is available at: <https://drive.google.com/file/d/0B2zMAIgfHF40RWE5jX3IsTFdsY0k/view?usp=sharing>.