Diversification of a Protein Kinase Cascade: IME-2 Is Involved in Nonself Recognition and Programmed Cell Death in *Neurospora crassa*

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ABSTRACT Kinase cascades and the modification of proteins by phosphorylation are major mechanisms for cell signaling and communication, and evolution of these signaling pathways can contribute to new developmental or environmental response pathways. The *Saccharomyces cerevisiae* kinase Ime2 has been well characterized for its role in meiosis. However, recent studies have revealed alternative functions for Ime2 in both *S. cerevisiae* and other fungi. In the filamentous fungus *Neurospora crassa*, the *IME2* homolog (*ime-2*) is not required for meiosis. Here we determine that *ime-2* interacts genetically with a transcription factor *vib-1* during nonself recognition and programmed cell death (PCD). Mutations in *vib-1* ($\Delta vib-1$) suppress PCD due to nonself recognition events; however, a $\Delta vib-1 \Delta ime-2$ mutant restored wild-type levels of cell death. A role for *ime-2* in the post-translational processing and localization of a mitochondrial matrix protein was identified, which may implicate mitochondria in *N. crassa* nonself recognition and PCD. Further, $\Delta vib-1$ strains do not produce extracellular proteases, but protease secretion reverted to near wild-type levels in a $\Delta vib-1 \Delta ime-2$ strain. Mass spectrometry analysis revealed that the VIB-1 protein is phosphorylated at several sites, including a site that matches the IME-2 consensus. The genetic and biochemical data for *ime-2* and *vib-1* indicate that IME-2 is a negative regulator of VIB-1 and suggest parallel negative regulation by IME-2 of a cell death pathway in *N. crassa* and provides insight into the evolution of kinases and their regulatory targets.

CELL-CELL signaling via kinase cascades is an essential mechanism for communication within and between organisms. Protein kinases are one of the largest protein families in eukaryotes and as much as 30% of any given eukaryotic proteome is phosphorylated (Deshmukh *et al.* 2010; Moses and Landry 2010). Although kinases have constrained target specificities, these proteins are often structured in a modular way, such that they can evolve new functions via interaction with scaffolds, adapters, or docking proteins (Bhattacharyya *et al.* 2006). In addition, duplication of kinase targets can

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result in reciprocal loss of phosphorylation sites and subfunctionalization of these targets and/or a gain of new phosphorylation sites, resulting in neo-functionalization (Amoutzias *et al.* 2010). Thus, changes in kinase structure, along with target duplication and divergence, can affect the structure and signaling output of kinase pathways. Although the major classes of kinases are conserved across fungal species, there is evidence for duplication, family expansion, and differences in domain organization, suggesting that fungi can change their kinase signaling pathways to accommodate changes in environment or developmental processes (Kosti *et al.* 2010).

In *Saccharomyces cerevisiae*, Ime2 is a serine/threonine protein kinase involved in the induction of meiosis and sporulation (Smith and Mitchell 1989). Ime2 has both early and late roles in meiosis, including the initiation of meiosis, meiotic DNA replication, meiotic divisions I and II, and spore formation (Benjamin *et al.* 2003; Honigberg 2004; Brush *et al.* 2012). Nutritional signals for meiosis converge at Ime1, a transcriptional regulator of Ime2, as well as at Ime2 itself, to coordinate meiotic initiation (Honigberg and Purnapatre

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Figure 1 Schematic for VIB-1 regulation of HI and cell death. VIB-1 is required for HI mediated by genetic differences at mating type, het-6, and het-c pin-c (Xiang and Glass 2002, 2004; Lafontaine and Smith 2012) in addition to activating PCD through additional unknown downstream effectors (Dementhon et al. 2006). het-6 incompatibility is mediated by un-24 (ribonucleotide reductase) and het-6 interactions (Micali and Smith 2006; Lafontaine and Smith 2012); mating-type incompatibility is mediated by mating type A-1, mating type a-1, and tol (Pittenger 1957; Newmeyer 1970; Glass et al. 1990; Shiu and Glass 1999); while het-c incompatibility is mediated by het-c pin-c interactions (Glass and Kaneko 2003; Kaneko et al. 2006); alternative HET-C polypeptides have been shown to physically interact (Sarkar et al. 2002). Phenotypic consequences of HI in N. crassa include growth inhibition and suppression of conidiation (Perkins 1988). Hyphal compartments within heterokaryons

that carry alternate *het* haplotypes (such as *het-c/pin-c*, *rnr/het-6*, or *mat-a/A/tol*) undergo compartmentation and rapid cell death (Glass and Kaneko 2003) and thus stain positive for vital dyes such as methylene blue (arrows).

2003; Kassir *et al.* 2003). One of the major roles of Ime2 is to contribute to activation of the major middle meiotic transcription factor *NDT80* (Pak and Segall 2002a; Sopko *et al.* 2002; Shin *et al.* 2010), in part through phosphorylation of the repressor Sum1 (Pak and Segall 2002b; Ahmed *et al.* 2009; Winter 2012). In addition, Ime2 directly phosphorylates Ndt80, which is associated with an increased ability to activate transcription of Ndt80 target genes (Sopko *et al.* 2002; Shubassi *et al.* 2003). Ndt80 binds to the middle sporulation element (MSE) and activates expression of middle meiotic genes (Chu and Herskowitz 1998; Chu *et al.* 1998); cells lacking Ndt80 arrest at pachytene, prior to nuclear division in meiosis I (Xu *et al.* 1995).

In the filamentous fungus *Neurospora crassa*, there is one IME2 homolog (ime-2), but three NDT80 homologs (female sexual development, fsd-1; vegetative incompatibility blocked, vib-1; and NCU04729) (Borkovich et al. 2004; Hutchison and Glass 2010). Homologs to IME1 and SUM1 are lacking in the N. crassa genome. Recently, we showed neither ime-2 nor the NDT80 homologs fsd-1, vib-1, or NCU04729 are involved in meiotic functions in N. crassa (Hutchison and Glass 2010). Mutations in ime-2 did not affect the transcription or activity of *fsd-1*, the homolog most closely related to S. cerevisiae NDT80. However, ime-2, vib-1, and fsd-1 mutants were affected in the production of female reproductive structures, termed protoperithecia. The development of protoperithecia in N. crassa is induced under conditions of nitrogen starvation (Westergaard and Mitchell 1947; Hirsh 1954). The $\Delta fsd-1$ and $\Delta vib-1$ mutants formed few protoperithecia under nitrogen starvation and an $\Delta fsd-1 \Delta vib-1$ mutant was female sterile. In contrast, a $\Delta ime-2$ strain produced protoperithecia under conditions where development of these structures is normally suppressed (nitrogen sufficiency) and significantly more protoperithecia are under nitrogen starvation conditions (Hutchison and Glass 2010). A deletion of *ime-2* restored protoperithecial development in a $\Delta fsd-1$ mutant, while a $\Delta ime-2 \Delta vib-1$ mutant showed a $\Delta vib-1$ phenotype (few protoperithecia). These observations indicate a network of regulatory interactions between ime-2 and the

NDT80 homologs *fsd-1* and *vib-1* during development of female reproductive structures in *N. crassa*.

The vib-1 mutant was first identified in a search for mutations that alleviate heterokaryon incompatibility (HI) in N. crassa (Xiang and Glass 2002). Heterokaryon incompatibility mediates nonself recognition and is a ubiquitous phenomenon in filamentous fungi (Aanen et al. 2010; Choi et al. 2012). Within a single filamentous fungal colony, individual hyphae can fuse and form interconnected networks (Fleissner et al. 2008; Read et al. 2010). However, if fusion occurs between strains that contain alternative specificities at heterokaryon incompatibility (het) loci, either the fusion cell is walled off and rapidly killed or the growth of the heterokaryon is inhibited (Glass and Kaneko 2003; Aanen et al. 2010) (Figure 1). In N. crassa, there are 11 het loci and genetic differences at any one of these 11 loci are sufficient to restrict heterokaryon formation (Glass and Dementhon 2006). Strains carrying genetic differences at het loci, but also loss-of-function mutations in vib-1, will form vigorous heterokaryons and do not show HI-associated programmed cell death (PCD) (Xiang and Glass 2002, 2004; Lafontaine and Smith 2012). vib-1 is also necessary for the expression of several genes known to be involved in cell death due to HI (Figure 1) (Dementhon et al. 2006). In Aspergillus nidulans, deletion of a vib-1 homolog, xprG, prevents secretion of extracellular proteases upon nitrogen or carbon starvation (Katz et al. 2006) and vib-1 mutants exhibit this phenotype as well.

In this study, we show that in *N. crassa*, a deletion of *ime-2* restores HI-induced PCD in a $\Delta vib-1$ strain and also restores the production of extracellular proteases. We further investigated the *ime-2* phenotype using transcriptional profiling to assess physiological differences between *ime-2* mutants and a wild-type strain and identified a possible role for IME-2 in mitochondrial homeostasis. Our data suggest that IME-2 is a negative regulator of a cell death pathway that functions in parallel to the VIB-1 HI pathway to specifically regulate nonself recognition and cell death when strains carry incompatible specificities at *het* loci.

Table 1 List of strains used in this study

Strain name	Genotype ^a	Origin or reference ^b
FGSC 2489	A	FGSC
FGSC 11308, FGSC 11309	Δvib -1::hph a; Δvib -1::hph A	FGSC
FGSC 17936, FGSC 17937	Δ ime-2::hph a; Δ ime-2::hph A	FGSC
FGSC 4564	ad-3B cyh-1 a ^{m1}	FGSC
R15-7	his-3; a	Dementhon <i>et al.</i> (2006)
C9-15	het-c2 pin-c2 thr-2 A	Smith <i>et al.</i> (2000)
C9-2	het-c2 pin-c2 thr-2 a	Smith <i>et al.</i> (2000)
Xa-3	het-c2 pin-c2 arg-5; pan-2 A	Xiang and Glass (2002)
IH3	het-c2 pin-c2 arg-5; a	$(9-2 \times Xa-3)$
5110		Dementhon <i>et al.</i> (2006)
R14-42	his-3 rid-1 Asad-1::hph A	Gift from P. K. T. Shiu: Rasmussen et al. (2008)
KD02-10	his-3: pvr-4: pan-2 a	Dementhon et al. (2006)
DVI.4	Δime-2::hph: Δvib-1::hph A	Hutchison and Glass (2010)
D49.10	his-3: ΔΝCU09915::hph: ΔΝCU04729::hph A	Hutchison and Glass (2010)
KD13-21	his-3: Δvib-1::hph A	Dementhon et al. (2006)
KD13-51	$\Delta vib-1::hph: pan-2 A$	Dementhon et al. (2006)
KD13-33	$\Delta vib-1::hph: pan-2 a$	Dementhon et al. (2006)
KD13-01	het-c2 pin-c2 thr-2: Δvib-1::hph a	Dementhon $et al.$ (2006)
KD13-23	his-3: het-c2 pin-c2: Δvib-1::hph: pan-2 A	Dementhon <i>et al.</i> (2006)
DVI.HIS.40	his-3: Aime-2: Avib-1 a	$DVI.4 \times KD02-10$
SV1	his-3:: $pccq1-vib-1-qfp: pvr-4: \Delta vib-1: pan-2 A$	Gift from J. Sun. Glass laboratory
BH13c	his-3::pcca1-vib-1-afp: $\Delta vib-1::hph$	$SV1 \times DVLHIS.40$
DI.PYR.4	$\Delta ime-2$::hph pvr-4 a	$DVI.4 \times KD02-10$
DI.HIS.10	his-3: Δime-2::hph a	$DVI.4 \times KD02-10$
DI.A.22	Δime-2::hph arg-5 het-c2 pin-c2 a	DVI.A.78 × D49.10
DVI.PYR.63	Δ ime-2::hph pvr-4: Δ vib-1::hph a	$DVI.4 \times KD02-10$
DVI.HIS.48	his-3: Δime-2::hph: Δvib-1::hph a	$DVI.4 \times KD02-10$
DVI.A.101	Δ ime-2::hph arg-5: Δ vib-1::hph a	$DVI.4 \times JH3$
DV.80	het-c2 pin-c2 arg-5: $\Delta vib-1 A$	$KD02-10 \times DVI.4$
R14-42arg4gfp	his-3::pccg1-arg-4-gfp rid-1 Δ sad-1::hph A	R14-42 [pccg1-arg-4-gfp]
Δ vib-1arg4gfp	his-3::pccg1-arg4-gfp; Δ vib-1::hph A	KD13-21 [pccg1-arg-4-gfp]
Δ ime2arg4gfp	his-3::pccg1-arg4-gfp; Δ ime-2::hph a	DI.HIS.10 [pccg1-arg-4-gfp]
2XKOarg4gfp	his-3::pccg1-arg4-gfp; Δ ime-2; Δ vib-1::hph	DVI.HIS.48 [pccg1-arg-4-gfp]
DVI.A.78	Δ ime-2::hph arg-5; Δ vib-1::hph a	$DVI.4 \times JH3$
D49VI.HIS.1	his-3; Δime-2::hph; ΔNCU09915::hph; Δvib-1::hph; ΔNCU04729::hph a	D49.10 × DVI.A.78
1XA	his-3::pccq1-vib-1(S60A)-qfp; het-c2 pin-c2 Δ vib-1::hph; pan-2 A	KD13-23 [pccg1- <i>vib-1</i> 1 \times PMA-gfp]
5XA	his-3::pccg1-vib-1(S60A, S413A, S537A, S542A, S545A)-gfp; het-c2 pin-c2 Δvib-1::hph; pan-2 A	KD13-23 [pccg1-vib-14 × PMA-gfp]
1XD	his-3::pccg1-vib-1(S60D)-gfp; het-c2 pin-c2 Δ vib-1::hph; pan-2 A	KD13-23 [pccg1- <i>vib-1</i> 1 × PMD-qfp]
5XD	his-3::pccg1-vib-1(S60D, S413D, S537D, S542D, S545D)-gfp; het-c2 pin-c2 Δvib-1::hph; pan-2 A	KD13-23 [pccg1-vib-14 × PMD-gfp]

^a Strains are of het-c1 pin-c1 genotype unless otherwise indicated.

^b "x", strains derived from crosses.

Materials and Methods

Strains and growth conditions

All strains used in this study are listed in Table 1. Deletion strains (FGSC 11308, FGSC 11309, FGSC 17936, and FGSC 17937) were constructed by the *Neurospora* program project grant (Colot *et al.* 2006) and obtained from the Fungal Genetics Stock Center (FGSC) (McCluskey 2003). Strains were grown on Vogel's minimal media (MM) (Vogel 1956) unless otherwise specified, and crosses were performed on Westergaard's media (Westergaard and Mitchell 1947). Transformations were performed as previously described (Margolin *et al.* 1997). To obtain forced heterokaryons, conidial sus-

pensions from strains of complementary auxotrophic markers were mixed and plated on minimal media. Growth rates were determined by growing strains in race tubes. Protoperithecial development was assessed over a 9-day time period of growth on water agar (Hutchison and Glass 2010).

Protease and cell death assays

For extracellular protease assays, strains were grown (in triplicate) in MM (Vogel 1956) overnight. Protease assays were performed as described previously (Dementhon *et al.* 2006). Cell death frequency was measured by staining with methylene blue (Hutchison *et al.* 2009). Heterokaryons were inoculated onto MM overlaid with cellophane, grown for

2–3 days, and stained for 1–2 min with 0.003% methylene blue. Approximately 20 random images were taken and the percentage of dead (blue) hyphal compartments was determined.

RNA extraction and quantitative RT-PCR

RNA extraction was performed on mycelia ground in liquid nitrogen or on sections of mycelia grown on cellophane. Mycelia were mixed with 0.3 g of 0.5-mm silica beads and 1 mL of TRIzol (Invitrogen, Carlsbad, CA) and disrupted using a bead beater (Mini-BeadBeater-8; Biospec Products). RNA was extracted according to the manufacturer's protocol for TRIzol (Invitrogen). Samples were purified using an RNAeasy kit (QIAGEN, Valencia, CA) and DNA was removed with QIAGEN DNase (no. 79254) or Ambion Turbo DNase (no. AM2238). RNA concentration and quality were assessed using a Nanodrop (Thermo Scientific) and gel electrophoresis. Quantitative RT-PCR (Q-RT-PCR) was performed using an EXPRESS One-Step SYBR GreenER kit (Invitrogen) according to the manufacturer's protocol, run on an ABI 7300 machine, and analyzed with ABI 7300 system software. Actin mRNA was used as the endogenous control, and reactions were performed in triplicate.

Microarray analysis

Microarray slide production, hybridization, and analysis were performed as described in Tian et al. (2007). Neurospora microarray slides are available from the FGSC (http://www. fgsc.net/). Approximately 10 µg of DNase-treated RNA was used as a template for cDNA synthesis (ChipShot Indirect cDNA Synthesis kit; Promega, Madison, WI), and hybridizations were performed using ProntoPlus kits (Promega), according to manufacturer instructions. Slides were scanned using an Axon GenePix 4000B scanner and analyzed using GenePix Pro 6 software (Molecular Devices). Three independent hybridizations pooled from three biological replicates were performed. Data were analyzed using Bayesian analysis of gene expression levels (BAGEL) (Townsend and Hartl 2002). Microarray data were verified by Q-RT-PCR, using template RNA from an independent experiment. All microarray data were deposited at the Filamentous Fungal Gene Expression Database (Zhang and Townsend 2010) and the Gene Expression Omnibus (GEO) database (ID GSE35905). Functional category enrichment analysis was carried out through the Munich Information Center for Protein Sequences (MIPS) database (http://www.helmholtz-muenchen.de/ en/mips/projects/funcat) (Ruepp et al. 2004), which uses a hypergeometric distribution to calculate P-values.

Mitochondrial staining

Mitochondria were visualized using 10 μ M MitoTracker Red FM (Invitrogen; no. M22425) (Hickey *et al.* 2004). Approximately 10⁵ conidia were inoculated into a 30-ml flask of MM and shaken at 30° for ~6 hr. MitoTracker Red FM was added to 1-ml aliquots of the conidia, followed by shaking at 30° for an additional 15–20 min. Conidia were pelleted by

centrifugation and washed once with MM. Conidia were then spread on a MM plate and incubated at 30° for 5–10 min. Mitochondria were imaged using a Deltavision Spectris DV4 deconvolution microscope (Applied Precision Instruments). A stack of ~20 images was taken 0.2 μ m apart, deconvolved using SVI Huygens, and visualized using Bitplane Imaris software.

Protein extraction, immunoprecipitation, and Western blotting

Protein was extracted from mycelia for immunoprecipitation (IP), using a method adapted from the FGSC Neurospora protocol page (http://www.fgsc.net/neurosporaprotocols/ Immunoprecipitation%20final.pdf). Briefly, 20-30 g of mycelia was ground in liquid nitrogen and homogenized using a 6770 Freezer/Mill from the SPEX CertiPrep Group, using three cycles of 1 min precool, 1 min run time, and 1 min cool time, at a speed of 15 cycles per second (CPS). Homogenized mycelia were added to HEPES IP extraction buffer [50 mM HEPES (pH 7.4), 137 mM NaCl, 10% glycerol] containing complete mini-EDTA-free protease inhibitor and PhosSTOP phosphatase inhibitor (Roche) and vortexed to homogenization. Samples were centrifuged at 3400 rpm for 10 min. Supernatants were concentrated via centrifugation, using Vivaspin 15R protein concentrators [10,000 molecular weight cut off (MWCO); Sartorius Stedium Biotech]. Four milliliters of each sample was immunoprecipitated using Protein G Dynabeads (Invitrogen), according to manufacturer's instructions, with the following exceptions: mouse anti-GFP antibody (Roche) was incubated with the beads for 1 hr, and the sample was immunoprecipitated for 2 hr at 4°. Protein was removed from the beads by boiling for 5 min, and samples were run on a 4-15% Criterion Tris-HCl gel (Bio-Rad, Hercules, CA). Protein gels were either subjected to Western blot analysis using standard methods or stained with SimplyBlue SafeStain Coomassie G-250 stain (Invitrogen) to visualize protein, and gel bands of interest were extracted using a razor blade.

Mass spectrometry

Gel bands of interest from Coomassie-stained gels were cut out and minced into <1-mm² pieces. Protein bands were extracted following a protocol adapted from the University of California (Berkeley) QB3 Proteomics/Mass Spectrometry Laboratory (http://qb3.berkeley.edu/pmsl/protocols.htm). Briefly, gel pieces were washed in 500 μ l NH₄HCO₃ for 20 min. After discarding the NH₄HCO₃ wash, 150 µl of NH₄HCO₃ (100 mM) and 10 µl of DTT (45 mM) were added and samples were incubated for 15 min at 50°. After cooling to room temperature, 10 µl of iodoacetamide (100 mM) was added and samples were incubated at room temperature for 15 min in the dark. The solvent was discarded and gel pieces were washed in 500 µl of a 1:1 mixture of acetonitrile and NH₄HCO₃ (100 mM) for 20 min. Then, gel pieces were incubated for 10-15 min in 50 µl of acetonitrile and dried in a speedvac. Gel pieces were rehydrated in 10 μ l of NH₄HCO₃



Figure 2 A deletion of *ime-2* partially restores HI in $\Delta vib-1$ mutants. Heterokaryons of identical *het-c pin-c* haplotype (C, compatible) show a growth and conidiation pattern indistinguishable from that of homokaryotic strains, including WT (A) (FGSC 4564 + R15-7), $\Delta vib-1$ (C) (KD13-21 + KD13-51), $\Delta ime-2$ (E) (DI.PYR.4 + DI.HIS.10), and $\Delta vib-1 \Delta ime-2$ (G) (DVI. HIS.48 + DVI.PYR.63) heterokaryons (Table 1). Strains carrying incompatible *het-c pin-c* haplotypes (Inc, incompatible) in a WT background show growth inhibition and suppression of conidiation (B) (FGSC 4564 + C9-15). Heterokaryons of incompatible *het-c pin-c* haplotypes, but carrying homozygous $\Delta vib-1$ mutations are similar in phenotype to compatible $\Delta vib-1$ heterokaryons (D) (KD13-33 + KD13-1). Heterokaryons carrying incompatible *het-c pin-c* haplotypes and homozygous $\Delta ime-2$ mutations show

(25 mM) and digested overnight with either trypsin or endoproteinase C, according to the manufacturer's instructions. Proteins were extracted twice with a mix of 60% acetonitrile and 0.1% formic acid and once with 100% acetonitrile. Extracted proteins were dried using a speedvac and twodimensional mass spectrometry analysis was performed at the QB3 Proteomics/Mass Spectrometry facility.

Results

A deletion of ime-2 restores HI-mediated cell death in a $\Delta \text{vib-1}$ mutant

In N. crassa, if two individuals are genetically identical at all nonself recognition loci (termed het), they can undergo cell fusion to form a "compatible" heterokaryon that looks identical to a homokaryotic wild-type strain. However, if individuals are genetically different at any 1 of 11 het loci, hyphal fusion results in compartmentalization of the fusion cell and subsequent death; these individuals are referred to as "incompatible" (Glass and Kaneko 2003) (Figure 1). In addition to cell death, incompatible heterokaryons have a reduced growth rate and do not conidiate. Strains that contain loss-of-function mutations in vib-1 form vigorous heterokaryons even if they differ in het allelic specificity (Xiang and Glass 2002, 2004) (Figure 2, C and D). Whereas compatible wild-type heterokaryons conidiate upon contact with plate edges, Δvib -1 heterokaryons of identical or alternate *het-c pin-c* haplotype display deregulated conidiation (Figure 2, C and D).

Due to genetic interactions identified between *ime-2* and *vib-1* with regard to formation of female reproductive structures (Hutchison and Glass 2010), we evaluated whether IME-2 plays a role in HI by assessing the incompatibility phenotype of $\Delta ime-2$ and $\Delta vib-1 \Delta ime-2$ mutants compared to wild-type and $\Delta vib-1$ mutants. To evaluate HI, we used a forced heterokaryon approach, using strains with the same *het-c pin-c* specificity (*het-c1 pin-c1* and thus compatible) *vs.* using strains containing alternate *het-c pin-c* haplotypes (*het-c1 pin-c1* + *het-c2 pin-c2* and thus incompatible) (Kaneko *et al.* 2006; Hall *et al.* 2010) (Figure 1 and Table 1). Further, these strains have different, complementary auxotrophic markers; when isolates carrying different auxotrophic markers are placed on minimal media, only the heterokaryotic strain is able to grow.

Compatible heterokaryons of identical *het-c pin-c* haplotype of wild-type, Δvib -1, Δime -2, or Δvib -1 Δime -2 strains were phenotypically identical to a homokaryotic wild-type or mutant strain by itself (Figure 2, A, C, E, and G). Wildtype heterokaryons with incompatible combinations of *het-c pin-c* haplotypes exhibited a severely decreased growth rate

typical HI (F) (DI.HIS.10 + DI.A.22), while heterokaryons carrying incompatible *het-c pin-c* haplotypes and homozygous for $\Delta vib-1 \Delta ime-2$ mutations show partial restoration of HI (decreased growth rate and suppression of conidiation) (H) (DVI.PYR.63 + DVI.A.101).



Figure 3 A deletion of ime-2 restores wild-type levels of cell death in Δvib -1 incompatible heterokaryons. (A) Heterokaryons of compatible and incompatible het-c pin-c haplotype from Figure 2 were plated on minimal media overlaid with cellophane and grown for 1-3 days. The percentage of dead cell compartments was evaluated using microscopy and methylene blue staining (Xiang and Glass 2002; Hutchison et al. 2009). Heterokaryons of incompatible het-c pin-c haplotype but carrying homozygous vib-1 deletions were suppressed for cell death (Δvib -1), while the addition of $\Delta ime-2$ mutation in these strains restored wild-type levels of cell death ($\Delta vib-1 \Delta ime-2$). (B) Expression of ime-2, vib-1, and two HET domain genes (pin-c, tol) was assessed using quantitative RT-PCR. Deletion of vib-1 abolishes expression of HET domain genes during vegetative growth. Expression of pin-c and tol was not restored in a $\Delta vib-1 \Delta ime-2$ mutant.

and lack of conidiation (compare Figure 2B with 2A). Heterokaryons carrying incompatible *het-c pin-c* haplotype and homozygous deletions of ime-2 also showed an identical incompatible phenotype to wild type (compare Figure 2F with 2B), indicating that ime-2 is not required for HI. In contrast, homozygous Δvib -1 heterokaryons of incompatible het-c pin-c haplotype showed a phenotype more similar to compatible $\Delta vib-1$ heterokaryons (compare Figure 2D to 2C); a deletion of vib-1 suppresses HI. However, a heterokaryon carrying incompatible *het-c pin-c* haplotypes, plus homozygous $\Delta vib-1$ $\Delta ime-2$ mutations showed significantly poorer growth and conidiation than a $\Delta vib-1$ suppressed incompatible heterokaryon (compare Figure 2H to 2D). The $\Delta vib-1 \Delta ime-2$ het-c *pin-c* incompatible heterokaryon had a growth rate of 2.80 \pm 0.55 cm/day (Figure 2H), compared to a wild-type incompatible heterokaryon (1.16 \pm 0.53 cm/day; Figure 2B) and a $\Delta vib-1$ incompatible heterokaryon (6.7 \pm 0.40 cm/day; Figure 2D). In addition, the $\Delta vib-1 \Delta ime-2$ het-c pin-c incompatible heterokaryon produced fewer conidia compared to a $\Delta vib-1$ suppressed incompatible heterokaryon (compare Figure 2H to 2D) and exhibited an altered, dense growth pattern that did not resemble either single mutants (Δvib -1 or Δime -2; Figure 2, C and E) or the Δvib -1 Δime -2 compatible strain (Figure 2G). Thus, in a $\Delta vib-1$ mutant, a deletion of ime-2 partially restored the inhibited growth

and aconidial phenotype associated with nonself recognition and HI.

In addition to growth inhibition and absence of conidiation, wild-type heterokaryons carrying incompatible *het-c pin-c* haplotypes show substantial PCD, with \sim 30% of hyphal segments showing compartmentalization and death, which is assessed by staining with vital dyes such as methylene blue (Figure 1) (Glass and Kaneko 2003). The Δvib -1, $\Delta ime-2$, and $\Delta vib-1 \Delta ime-2$ heterokaryons of identical het-c *pin-c* haplotype exhibited very little cell death, $\sim 2-5\%$, which was similar to that of a wild-type compatible heterokaryon (Figure 3A). Wild-type heterokaryons carrying incompatible het-c pin-c haplotypes showed ~30% hyphal compartmentation and death (Xiang and Glass 2002). Similar to wild-type incompatible heterokaryons and consistent with the HI phenotype (Figure 2F), the $\Delta ime-2$ heterokaryon carrying incompatible *het-c pin-c* haplotypes showed $\sim 27\%$ hyphal compartmentation and death (Figure 3A). As observed previously (Xiang and Glass 2002), $\Delta vib-1$ heterokaryons carrying incompatible *het-c pin-c* haplotypes showed a substantially reduced cell death frequency that was similar to that of either $\Delta vib-1$ or wild-type compatible heterokaryons (~2–5%). In contrast, the $\Delta vib-1 \Delta ime-2$ heterokaryon carrying incompatible het-c pin-c haplotypes showed a cell death frequency similar to that of wild-type incompatible heterokaryons ($\sim 25\%$)

(Figure 3A). These data indicate that loss-of-function mutations in *ime-2* restored hyphal compartmentation and death to $\Delta vib-1$ heterokaryons carrying incompatible *het-c pin-c* haplotypes.

Many proteins involved in HI, like PIN-C, contain a conserved protein domain of unknown function, termed HET (PF06985) (Espagne et al. 2002). HET domains are filamentous fungal-specific protein domains that can cause an HI-like cell death when overexpressed (Paoletti and Clave 2007). Previously, it was shown that *vib-1* is necessary for the expression of HET domain genes pin-c, tol, and het-6, which are required for HI in N. crassa (Dementhon et al. 2006) (Figure 1). To test whether restoration of cell death in $\Delta vib-1 \Delta ime-2$ strains correlated with HET domain gene expression, we performed quantitative RT-PCR for pin-c and tol in wild-type, Δvib -1, Δime -2, and Δvib -1 Δime -2 strains (Figure 3B). As expected, expression of *pin-c* and *tol* was not detected in a $\Delta vib-1$ mutant. Although expression of *pin-c* and tol in $\Delta ime-2$ strains was not significantly different from that in wild type, expression of vib-1 was significantly increased, suggesting that IME-2 negatively regulates vib-1 expression levels. In the $\Delta vib-1 \Delta ime-2$ strain, however, expression of pin-c and tol was similar to expression levels observed in the Δvib -1 strain (Figure 3B), indicating that restoration of cell death by $\Delta ime-2$ mutations in a $\Delta vib-1$ strain carrying incompatible het-c pin-c haplotypes is not due to induced expression of pin-c.

Microarray analysis reveals that mutations in ime-2 affect mitochondrial function

Because we identified a genetic interaction between vib-1 and ime-2 during HI, we evaluated what physiological processes were affected in the $\Delta ime-2$ mutant by performing gene expression profiling of wild type vs. a $\Delta ime-2$ strain under nitrogen starvation conditions. The initiation of female reproductive structures (protoperithecia) in N. crassa is regulated by the availability of nitrogen (Westergaard and Mitchell 1947; Hirsh 1954), and we hypothesized that differences in gene expression between wild-type and $\Delta ime-2$ deletion strains may be more pronounced under nitrogen starvation conditions (Hutchison and Glass 2010). The wildtype strain (FGSC 2489) and a $\Delta ime-2$ strain (FGSC 17937) were grown overnight (~16 hr) in minimal media. The mycelia were then washed and subsequently transferred to a flask containing minimal media without nitrogen and grown for an additional 4 hr (see Materials and Methods). Mycelia from both strains were harvested for RNA extraction and microarray analysis. Three replicate microarrays, including dye swaps, were performed.

In the $\Delta ime-2$ strain, a total of 187 genes showed a significant decrease in expression level of at least 1.5-fold. Functional category analysis (Ruepp *et al.* 2004) of this gene set showed enrichment for energy (P < 0.0001); transcription (P < 0.05); cellular transport (P < 0.005); and transposable elements, viral and plasmid proteins (P < 1e-11) (Figure 4). Many of the genes in the energy functional category belonged

to pathways involving electron transport and respiration. Furthermore, the enrichment of genes belonging to the "transposable elements, viral and plasmid proteins" functional category was due almost exclusively to genes belonging to the mitochondrial genome. In fact, 13 of the 29 genes composing the mitochondrial genome showed decreased expression levels in the $\Delta ime-2$ strain compared to wild type (Supporting Information, Table S1), a significant enrichment (P < 0.006). These observations suggest mitochondrial impairment in the $\Delta ime-2$ mutant and are consistent with the fact that $\Delta ime-2$ strains exhibit a slower growth rate (3.5–4 cm/day) compared to wild-type strains (~7 cm/day).

A total of 506 genes showed increased expression levels in the $\Delta ime-2$ strain compared to wild type (Figure 4). These genes were enriched in a variety of functional categories (Ruepp *et al.* 2004), including metabolism (P < 1e-11); energy (P < 1e-17); cell-cycle and DNA processing (P < 1e-17); 0.0001); transcription (P < 0.005); protein synthesis (P <1e-24); protein fate (P < 1e-5); protein with binding function or cofactor requirement (P < 1e-44); regulation of metabolism and protein function (P < 0.0005); cellular transport (P < 1e-14); cellular communication and signal transduction (P < 1e-4); cell rescue, defense, and virulence (P < 0.0005); interaction with the environment (P < 1e-6); cell fate (P < 1e-8); development (P < 0.005); diogenesis of cellular components (P < 1e-12); and cell-type differentiation (P < 1e-6). Although we hypothesized that the constitutive production of protoperithecia in $\Delta ime-2$ strains (Hutchison and Glass 2010) may be due to a defect in nitrogen sensing, we did not observe significant differences in gene expression with respect to nitrogen metabolism genes or genes involved in the metabolism of amino acids. Thus, the array data instead suggest that $\Delta ime-2$ mutants are not deficient in nitrogen sensing specifically, but that these strains may have a more general nutrient-sensing defect.

Interestingly, genes involved in mitochondrial biogenesis (within the biogenesis of cellular components category) were also significantly enriched among genes that showed increased expression levels in the $\Delta ime-2$ mutant (P < 0.0001). A recent study by Keeping et al. (2010) used mass spectrometry as well as computational methods to compile a comprehensive list of 738 genes that compose the N. crassa mitochondrial proteome. Using this data set, we asked whether nuclear-encoded mitochondrial genes were expressed differently between wild-type and $\Delta ime-2$ strains. In fact, these genes were significantly enriched (P < 1e-18) in the set of 506 genes that showed increased expression levels in the $\Delta ime-2$ mutant. As previously mentioned, 13 of the 29 genes composing the mitochondrial genome showed lower expression levels in the $\Delta ime-2$ strain (Table S1). These data suggest that the $\Delta ime-2$ mutant may have impaired mitochondria (evidenced by decreased expression of genes within the mitochondrial genome), resulting in a regulatory feedback loop such that $\Delta ime-2$ strains increase expression of nuclear-encoded mitochondrial genes to compensate for this defect. Overall, the microarray data suggest



Figure 4 Functional category analysis of gene expression differences in wild type vs. the $\Delta ime-2$ mutant. Shown is the distribution of significantly enriched MIPS functional categories (http://www.helmholtz-muenchen.de/en/mips/projects/funcat) (Ruepp *et al.* 2004) for the microarray data set of wild type compared to a $\Delta ime-2$ deletion strain. A total of 187 genes showed a reduction in expression in $\Delta ime-2$ relative to wild type, while 506 genes showed an increase in expression level in the $\Delta ime-2$ strain relative to wild type.

that IME-2 plays a role in mitochondrial function in *N. crassa*. We therefore assessed, via microscopy, whether mitochondrial morphology or protein localization was altered in $\Delta ime-2$ mutants compared to wild type and the $\Delta vib-1$ mutants.

ime-2 mutants affect post-translational processing of the mitochondrial protein ARG-4

To evaluate the mitochondrial phenotype in wild-type, Δvib -1, $\Delta ime-2$, and $\Delta vib-1 \Delta ime-2$ strains, we stained conidial germlings with MitoTracker Red FM and also transformed each strain with the nuclear-encoded GFP-tagged mitochondrial marker gene encoding ARG-4 (Bowman et al. 2009). The nuclear arg-4 locus encodes acetylornithine-glutamate acetyltransferase (arginine biosynthetic pathway), which is imported into the mitochondrial matrix (Cybis and Davis 1975). In wild-type hyphae, mitochondria appear as long tubules in the apical regions of hyphae and as more punctate structures farther back from the hyphal tip (Bowman et al. 2009). When stained with MitoTracker Red FM, all strains $(\Delta vib-1, \Delta ime-2, \text{ and } \Delta vib-1 \Delta ime-2)$ looked identical to wild type (Figure 5A) and had brightly stained, long, tubular mitochondrial networks. Similarly, when mitochondria were visualized via ARG-4-GFP localization, wild type and the $\Delta vib-1$ mutant exhibited long, tubular mitochondria that looked identical to the MitoTracker Red FM-stained mitochondria (Figure 5B). However, $\Delta ime-2$ strains exhibited an altered localization pattern. Instead of long, tubular structures, ARG-4-GFP localized either to vesicles or to punctate structures or was diffuse in the cytoplasm (Figure 5B). Mito-Tracker Red FM can permeate the cell membrane and accumulates in mitochondria based on membrane potential (Macho *et al.* 1996; Poot *et al.* 1996), and thus it is likely that the mitochondria observed in all strains, including the $\Delta ime-2$ strains, were active and have a functioning membrane potential. However, the lack of ARG-4-GFP localization in the $\Delta ime-2$ strain suggested that IME-2 may play a role in protein targeting to the mitochondria. Interestingly, the $\Delta vib-1 \Delta ime-2$ strain restored localization of ARG-4-GFP to tubular mitochondria (Figure 5B).

In yeast, the ARG-4 homolog Arg7 undergoes a posttranslational autoproteolytic processing step that results in the formation of two smaller subunits, each of which localizes to the mitochondrial matrix, where they associate in a complex (Abadjieva *et al.* 2000). The autoproteolytic activity of yeast Arg7 is dependent on a threonine residue, and this residue is conserved in *N. crassa* ARG-4. Therefore, we determined whether the defect in ARG-4-GFP localization in $\Delta ime-2$ strains was due to a defect in protein processing. Two distinct bands for *N. crassa* ARG-4-GFP (~75 kDa and 50 kDa) have previously been reported (Bowman *et al.* 2009), which is expected if ARG-4 is proteolytically cleaved at the conserved threonine residue. In the wild-type and $\Delta vib-1$ strains, two ARG-4-GFP bands at ~75 kDa and 50 kDa were detected (Figure 6, A and B). However, in $\Delta ime-2$ strains



Figure 5 Mutations in ime-2 affect localization of the mitochondrial protein ARG-4. (A) Mitochondria stained with MitoTracker Red FM in FGSC 2489 (WT), FGSC 11308 (Δ*vib-1*), FGSC 17937 (Δime -2), and DVI.4 (Δvib -1) $\Delta ime-2$) (Table 1). The right column is an enlargement of the region highlighted by a white box in the center column. (B) Localization of ARG-4-GFP to mitochondria in wild-type and deletion strains transformed with pccq1-arq-4gfp (Table 1). Localization of ARG-4-GFP to mitochondria in wild type is identical to mitochondria stained by MitoTracker Red FM and to that previously reported (Bowman et al. 2009). ARG-4-GFP localization in the Δvib -1 and the Δvib -1 $\Delta ime-2$ mutants was identical to that in WT. However, in the $\Delta ime-2$ strain, mitochondrial tubule structures were not observed and instead ARG-4-GFP localized to either vesicles or punctae or was diffuse in the cytoplasm. Bar, 5 µm.

much less ARG-4-GFP protein was present and only the 50-kDa ARG-4-GFP band was detected. Despite the lack of detectable full-length ARG-4 in the $\Delta ime-2$ mutants, these mutants were not arginine auxotrophs. In *S. cerevisiae, arg7* mutants exhibit a leaky Arg phenotype (Crabeel *et al.* 1997). Consistent with microscopy results, the $\Delta vib-1 \Delta ime-2$ strain showed a wild-type pattern for ARG-4 processing (Figure 6, A and B). In fact, $\Delta vib-1 \Delta ime-2$ strains appeared to produce more ARG-4-GFP than wild-type or $\Delta vib-1$ deletion strains.

The arg-4-gfp construct used to visualize ARG-4-GFP localization is under the regulation of the ccg-1 promoter (Bowman et al. 2009), which is commonly used for constitutive gene expression in N. crassa. The wild-type and the Δvib -1, Δime -2, and Δvib -1 Δime -2 strains transformed with the pccg-arg-4-gfp construct also contain a native copy of arg-4. Thus, we quantified the transcription of arg-4 in comparison to ccg-1, using Q-RT-PCR in all strains to assess expression levels of arg-4 (Figure 6D). Expression levels for arg-4 (a readout for both the ccg-1-regulated and the resident arg-4 genes) were very similar between wild type and $\Delta vib-1$ and $\Delta ime-2$ mutants. These data indicate that the differences observed in ARG-4 protein levels between wildtype and $\Delta ime-2$ strains were not due to decreased expression levels of *arg-4* in the $\Delta ime-2$ mutant (Figure 6, A, B, and D). However, the expression level of arg-4 was significantly elevated in the $\Delta vib-1 \Delta ime-2$ strain, consistent with increased ARG-4 protein levels in this strain detected via Western blot (Figure 6B). Wild type and the $\Delta ime-2$ mutant also showed similar levels of *ccg-1* expression, while the $\Delta vib-1$ and $\Delta vib-1 \Delta ime-2$ strains showed slightly lower levels of ccg-1 expression (Figure 6D). Because the ccg-1 promoter was not downregulated in the $\Delta ime-2$ strain or upregulated in the $\Delta vib-1 \Delta ime-2$ strain, it is likely that differences in proteins

levels of ARG-4 are not due to the regulation of the *arg-4* transgene. Either the increased *arg-4* transcription in the $\Delta vib-1 \Delta ime-2$ strain originated from the native *arg-4* locus or *arg-4* transcripts were stabilized in this strain. Thus, *ime-2* and *vib-1* affect the post-translational modification of ARG-4 and, to some degree in the $\Delta vib-1 \Delta ime-2$ strain, transcriptional regulation of *arg-4*.

Mutations in ime-2 revert the protease secretion phenotype of vib-1 mutants

Mutations in the transcription factor vib-1 cause a visible phenotype during vegetative growth, such that $\Delta vib-1$ mutants show pinkish (rather than orange) conidial pigmentation, deregulated conidiation, decreased aerial hyphae formation (Figures 2C and 7A), and a slight decrease in growth rate compared to wild type (Xiang and Glass 2002; Dementhon et al. 2006; Hutchison et al. 2009). In contrast, wild type and the $\Delta ime-2$ mutant show robust aerial hyphae formation and conidiation at the top of the tube or edge of the plate and were nearly indistinguishable, with the exception of the slightly yellow pigmentation observed in the $\Delta ime-2$ mutant (Figure 7A). In the $\Delta vib-1 \Delta ime-2$ mutant, aerial hyphae formation was restored and conidiation occurred only at the top of the slants, a phenotype similar to wild type and the $\Delta ime-2$ mutant (Figure 7A). However, conidia of the $\Delta ime-2$ $\Delta vib-1$ mutant are pinkish in color, like those of the $\Delta vib-1$ single mutants.

In addition to the vegetative conidiation phenotype, Δvib -1 mutants do not secrete extracellular proteases in response to nitrogen or carbon starvation (Dementhon *et al.* 2006), a phenotype similar to the *vib*-1 homolog in *A. nidulans*, *xprG* (Katz *et al.* 2006). We therefore evaluated whether mutations in *ime*-2 restored protease activity in the Δvib -1 mutant. When



Figure 6 Strains carrying a deletion in *ime-2* affect the post-transcriptional regulation of ARG-4. (A) Western blot for ARG-4-GFP in wild-type, Δvib -1, Δime -2, and Δvib -1 Δime -2 strains, with molecular weight ladder (kDa) on the left. ARG-4-GFP is detected as two distinct bands, ~75 kDa and 50 kDa in WT, as previously reported (Bowman *et al.* 2009). (B) Longer exposure of the blot from A, more clearly showing the 50-kDa ARG-4-GFP in the Δime -2 strain. (C) Western blot of β -tubulin showing that equal amounts of protein were loaded in each well. (D) Quantitative RT-PCR of *arg*-4 and ccg-1 transcript levels in wild-type, Δvib -1, Δime -2, and Δvib -1 Δime -2 strains.

nitrogen was provided, extracellular proteases were not induced in the wild-type, Δvib -1, Δime -2, or Δvib -1 Δime -2 strains (Figure 7B). When the wild-type strain was transferred to nitrogen starvation medium, extracellular protease activity was induced, while no activity was detected in the Δvib -1 mutant. A strain carrying a deletion of Δime -2 produced slightly elevated levels of proteases in response to nitrogen starvation (Figure 7, B and C). However, unlike the Δvib -1 mutant, the Δvib -1 Δime -2 mutant showed wildtype protease activity in response to nitrogen starvation (Figure 7, B and C), indicating that loss-of-function mutations in *ime*-2 suppressed the defect in protease secretion in Δvib -1 mutants.

Based on the Δvib -1 Δime -2 phenotype, we hypothesized that additional regulators may be functioning redundantly to *vib-1* to restore protease production. Two obvious candidate genes that may have redundant functions with vib-1 are the vib-1 paralogs fsd-1 and NCU04729. Previously, we determined that strains containing a deletion of NCU04729 were indistinguishable from wild type under all conditions, while $\Delta fsd-1$ mutants shows defects in protoperithecial formation and ascospore maturation (Hutchison and Glass 2010); neither fsd-1 nor NCU04729 affect cell death due to HI. We therefore tested the ability of a $\Delta vib-1 \Delta ime-2 \Delta fsd-1$ Δ NCU04729 deletion strain (Table 1, strain D49VI.HIS.1) to produce extracellular proteases. As shown in Figure 7, the quadruple mutant (Δvib -1 Δime -2 Δfsd -1 $\Delta NCU04729$) produced near wild-type levels of extracellular proteases (Figure 7B), indicating that neither fsd-1 or NCU04729 were responsible for the restoration of protease secretion in the $\Delta vib-1 \Delta ime-2$ mutant.

VIB-1 is phosphorylated at a predicted IME-2 consensus site

In S. cerevisiae, the consensus sequence for Ime2 phosphorylation (R-P-x-S/T-A/R-G) has been well characterized (Holt et al. 2007; Moore et al. 2007). We analyzed the N. crassa genome for matches to the yeast Ime2 phosphorylation consensus sequence, using the Scansite program (http://scansite. mit.edu/) (Obenauer et al. 2003) with a slightly modified phosphorylation consensus identified for N. crassa (R-P-x-S/ T-P/A/R-G) (L. Holt Laboratory, unpublished data). There are 30 total predicted phosphorylation targets of IME-2 in the N. crassa genome (Table S2). Consistent with the role of ime-2 in protoperithecial formation, one of the predicted IME-2 phosphorylation targets present in the Scansite data set is NIT-2 (NCU09068), a major regulator of nitrogen utilization. In addition, both AL-1 (albino-1; NCU00552), a phytoene dehydrogenase involved in carotenoid biosynthesis (Schmidhauser et al. 1990), and NRC-2 (NCU01797), a serine-threonine kinase involved in regulation of entry into the conidiation pathway and conidial development (Kothe and Free 1998), were present in the Scansite data set. $\Delta ime-2$ strains appear to have a slightly different conidiation phenotype from that of wild type (WT), including less pigmentation and fewer conidia (Figures 2E and 7A). The N. crassa homolog of the yeast protein kinase Ste20 (NCU03894) is also a predicted IME-2 phosphorylation target. Ste20 and its homologs in mammals (Mst1 and Mst2) have been previously shown to have a role in the apoptotic signaling cascade (Madeo et al. 2009; Radu and Chernoff 2009). Additionally, the VIB-1 protein contains a match for the Ime2 consensus, while neither of the other two NDT80 homologs in N. crassa (fsd-1 or NCU04729) have an Ime2 consensus site. From a phosphoproteomics study (A. Leeder and N. L. Glass, unpublished results), we identified a phosphopeptide for VIB-1 at the predicted IME-2 consensus site (RPRS*60), as well as four additional phosphorylation sites (MPQS*413, PSKS*537, and RHGS*⁵⁴²HGS*⁵⁴⁵) (Figure 8A).

To test whether the predicted IME-2 consensus site was necessary for VIB-1 function, we constructed mutant vib-1 alleles such that the IME-2 site was mutated to alanine (predicted to be phospho-null; S60-to-A mutation) (1XA, Table 1) or mutated to aspartate (phospho-mimetic; S60-to-D mutation) (1XD, Table 1). The growth rates of strains carrying the vib-1^{S60A} and vib-1^{S60D} alleles were identical to those of wild type as was nuclear localization of VIB-1^{S60A}-GFP and VIB-1^{S60D} -GFP (Figure S1 and Figure S2). Although identical in phenotype to wild type, both the vib-1^{S60A}-gfp and the vib-1^{S60D} -gfp strains showed slightly lower protease levels, suggesting that phosphorylation of the S60 site contributes to VIB-1 activity (Figure 7C). Similarly, mutations of the IME-2 consensus sequence in VIB-1 significantly reduced the numbers of protoperithecia produced in vib-1^{S60A}-gfp and vib-1^{S60D} -gfp strains under conditions of nitrogen starvation (Figure S3). These data indicate that mutations in the Ime2 consensus site on VIB-1 negatively affect protoperithecial development.



Figure 7 An ime-2 deletion restores wild-type conidiation patterns and protease production to a $\Delta vib-1$ mutant. (A) Wild-type (FGSC 2489), *Δvib-1* (FGSC 11308), *Δime-2* (FGSC 17936), and $\Delta vib-1 \Delta ime-2$ (DVI.4) strains grown on minimal media slants. (B) Extracellular protease activity of wild-type and deletion strains (from A), as well as that of the Δvib -1 Δime -2 Δfsd -1 $\Delta NCU04729$ strain (D49VI. HIS.1), was assessed on media with and without nitrogen. Only the vib-1 deletion strain showed inability to secrete proteases. Protease activity units are normalized to WT in nitrogen starvation (-nitrogen) media. (C) Extracellular protease activity of vib-1 phospho-mutants (vib-1^{560A} 1XA, vib-1560D 1XD, vib-1560A;5413A;5537A;5542A;5545A 5XA, and vib-1^{S60D;S413D;S537D;S542D;S545D} 5XD) (Table 1). Data are shown as the fold increase of protease secretion compared to that in the strains grown on media containing nitrogen. Asterisks (*) in B and C indicate strains with protease production significantly different from that in WT (P < 0.05).

To assess the role of the Ime2 phosphorylation sites on HI, the *vib-1*^{S60A}-*gfp* and *vib-1*^{S60D}-*gfp* alleles were transformed into Δvib -1 strains of *het*-*c*2 *pin*-*c*2 haplotype (Table 1). Each strain was then forced in a heterokaryon with a Δvib -1 strain of *het*-*c*1 *pin*-*c*1 haplotype. The Δvib -1 mutation is recessive,

such that a single functional copy of *vib-1* in a heterokaryon is sufficient to trigger HI (Xiang and Glass 2002; Dementhon *et al.* 2006) (Figure 8B, compare panels 3 and 4). When strains containing *vib-1*^{S60A}-gfp or *vib-1*^{S60D}-gfp were forced in a heterokaryon with a $\Delta vib-1$ strain of incompatible *het-c*



- a* 55 ...VRPR**S**R...62
- b 390 ...RVSSMDLPRTAFTFTSAKQMPQSPMQ...417
- c-e 533 ...PSKSPRHGSHGSLTNE...550





Figure 8 Phosphorylation of VIB-1 and phenotype of phospho-site mutants. (A) VIB-1 is phosphorylated on multiple sites (determined by mass spectrometry), including a predicted IME-2 consensus site. The asterisk indicates the phosphorylation site that matches the consensus for yeast Ime2. (B) Mutations of the IME-2 consensus phosphorylation site in VIB-1 do not affect growth inhibition and suppression of conidiation associated with HI. Δvib -1 compatible (1, KD13-21 + KD13-51) and incompatible (2, KD13-21 + DV.80) heterokaryons (suppressed for HI) are shown. A heterokaryon with only one functional copy of vib-1 and carrying incompatible *het-c pin-c* haplotypes shows typical HI (Δvib -1 mutations are recessive) (4, KD13-51 + JH1) compared to a heterokaryon carrying one functional copy of vib-1, but of compatible het-c pin-c haplotype (3, KD13-21 + FGSC 6103). Heterokaryons with a phospho-null mutation (S60 to A) (5, 1XA + KD13-21) or a phospho-mimetic mutation (S60 to D) (6, 1XD + KD13-21) at the predicted IME-2 consensus site in VIB-1 exhibited a typical wild-type HI phenotype. (C) Mutations of the IME-2 consensus phosphorylation site in VIB-1 do not affect cell-death percentages associated with HI. However, mutations at all five VIB-1 phosphorylation sites reduce cell-death percentages, although growth inhibition is still observed. All incompatible (Inc) heterokaryons exhibit significantly higher cell death than compatible (C) heterokaryons. Cell death percentages for incompatible heterokaryons labeled with an asterisk (*) were significantly different (P < 0.05) from those for heterokaryons labeled with an arrowhead $(\mathbf{\nabla})$.

pin-c haplotype, a HI response was triggered that was indistinguishable from a wild-type HI phenotype (Figure 8B, panels 5 and 6). In addition, the *vib-1*^{S60A}-*gfp* and *vib-1*^{S60D}- *gfp* incompatible heterokaryons displayed wild-type levels of cell death (Figure 8C).

We also constructed strains where all five identified phosphorylation sites were mutated to alanine (S60A, S413A, S537A, S542A, S545A, and 5XA) or aspartate (S60D, S413D, S537D, S542D, S545D, and 5XD) (Table 1). Both the 5XA and 5XD strains grew significantly slower than WT (Figure S2) and were also impaired in cell death during HI (Figure 8C). The 5XA and 5XD mutants also showed a significant decrease in protoperithecial formation compared to WT (Figure S2), with the 5XD strain showing the most significant reduction, with values similar to those of the Δvib -1 and Δvib -1 Δ ime-2 mutants. Consistent with our data that five amino acid substitutions in VIB-1 negatively affect its function, the 5XA and 5XD strains showed less protease activity than WT (P = 0.05), with values similar to those of the 1XA and 1XD strains.

Discussion

Diversification of kinase cascades may provide a mechanism for eukaryotes to evolve new developmental pathways or adapt to new environments (Bhattacharyya et al. 2006). In this study, we showed that N. crassa ime-2 regulates cell death due to HI in the absence of vib-1 and also regulates post-translational processing of the mitochondrial matrix protein ARG-4. IME2 homologs have not been previously implicated in programmed cell death, but recent studies in yeast and other fungi have provided evidence that Ime2 and its homologs can function in cellular processes other than meiosis. For instance, Strudwick et al. (2010) described a role for Ime2 in yeast pseudohyphal formation, and studies in other fungal species showed that although Ime2 homologs often function in sexual differentiation or nutrient sensing, they are not generally meiotic regulators (Irniger 2011). Our results provide additional evidence that the function of the Ime2 pathway differs among fungal species and implicate N. crassa ime-2 in nonself recognition and programmed cell death.

Data from this study as well as from a previous study (Hutchison and Glass 2010) indicate that ime-2 interacts genetically with the transcription factor vib-1. For some phenotypes, such as HI, protease production, and conidiation, *ime-2* is epistatic to *vib-1*. However, for other phenotypes, including protoperithecial formation, HET domain gene expression, and ARG-4 localization and protein processing, vib-1 is epistatic to *ime-2*. These data suggest that the *ime-2/vib-1* signaling pathway is not a simple, linear interaction, but that there are other genetic interactors present depending on the cellular process. We propose that the overall structure of the pathway (with the IME2 homolog functioning upstream of the NDT80 homolog) is likely conserved, that IME-2 negatively regulates VIB-1 (likely at the protein level), and that IME-2 regulates a parallel pathway that functions redundantly with VIB-1 (Figure 9) to regulate HI and protease production. In this scenario, protease production and cell



death in a Δvib -1 Δime -2 strain is restored. Mutation of components in the parallel cell-death induction pathway in addition to vib-1 mutations would ameliorate cell death and HI completely, regardless of the presence or absence of *ime*-2. Further experiments will be needed to identify IME-2 targets and additional members of this pathway.

Loss-of-function mutations in *vib-1* result in an inability to secrete proteases in response to nitrogen starvation and presence of extracellular protein, a phenotype that is suppressed in $\Delta ime-2 \Delta vib-1$ mutants. In eukaryotes, including *S. cerevisiae*, caspase proteases (metacaspases in *S. cerevisiae*) are integral in the activation of the apoptotic cell-death cascade (Madeo *et al.* 2004; Tait and Green 2010; Abdelwahid *et al.* 2011). In *N. crassa*, metacaspases are not required for HI-induced cell death (Hutchison *et al.* 2009). However, it is possible that a link between HI-induced cell death and VIB-1/IME-2– regulated proteases occurs in *N. crassa* by the utilization of nonmetacaspase proteases to induce cell death.

Mitochondria are key players in apoptosis and cell-death pathways (Tait and Green 2010). During apoptotic cell death, the pro-apoptotic Bcl-2 proteins BAX and BAK can cause the mitochondrial outer membrane to permeabilize, disrupting mitochondrial function, energy production, and redox potential and promoting the release of pro-apoptotic factors such as cytochrome c (Degterev and Yuan 2008; Tait and Green 2010; Abdelwahid et al. 2011). In fungi, mitochondria have also been implicated in apoptotic cell death and have important roles in life span and senescence (Madeo et al. 2004; Maheshwari and Navaraj 2008; Sharon et al. 2009). Mutations in ime-2 affected localization and post-transcriptional processing of ARG-4, a phenotype that was restored to a wild-type pattern in the $\Delta vib-1 \Delta ime-2$ mutant. The observation that IME-2 affects post-translational processing of a mitochondrial matrix protein suggests that the parallel cell-death pathway could be acting through the mitochondria or a mitochondria-related pathway. S. cerevisiae IME2 and its homolog in Schizosaccharomyces pombe, mde3/pit1, have a role in meiosis (Abe and Shimoda 2000; Kassir et al. 2003; Honigberg 2004) as well as in pseudohyphal growth

Figure 9 Model for the ime-2 and vib-1 genetic pathway. (A) VIB-1 positively regulates HI and cell death, as well as HET domain gene expression (via a separate mechanism). IME-2 negatively regulates additional HI cell death mediators that function in parallel to the VIB-1 pathway. Due to the role of IME-2 in the processing of a mitochondrial protein, we reasoned that the alternate cell-death effectors and/or pathway could be acting through the mitochondria. Further, since deletions in ime-2 do not exhibit increased cell death during HI, we suggest that VIB-1 negatively regulates these alternate death effectors. (B) VIB-1 also positively regulates secreted proteases and, similar to the mechanism for HI cell-death regulation, IME-2 negatively regulates a parallel pathway for secreted proteases. Deletions in ime-2 cause a significant increase in secreted proteases, and thus in this pathway, VIB-1 does not regulate the parallel pathway for secreted proteases. Further, IME-2 appears to negatively regulate VIB-1 with respect to protoperithecial development via a separate pathway. For both pathways, it is likely that IME-2 is regulated by cellular signals of nitrogen availability.

(Strudwick *et al.* 2010). Both of these developmental processes are associated with nitrogen starvation, suggesting an additional role for Ime2 in nutrient sensing, similar to that proposed for *N. crassa* (Hutchison and Glass 2010). It will be of interest to assess whether *ime2* Δ mutants in *S. cerevisiae* also have in common a defect in the localization and processing of Arg7 (ortholog of *N. crassa arg-4*), as observed in *N. crassa* Δ *ime-2* mutants. Further experiments on the relationship between *ime-2/IME2* and mitochondrial function are warranted.

We identified five phosphorylation sites on VIB-1, including a site that matches the predicted consensus site for Ime2. However, mutations in the Ime2 consensus site in VIB-1 that are predicted to result in phospho-null mutations (S to A) or activating mutations (S to D) resulted in no to only subtle phenotypic differences from WT. In particular, no role for the Ime2 phosphorylation site was observable for HI. Strains containing vib-1 alleles containing five S-to-A or five S-to-D mutations showed more severe defects, particularly reduced vegetative growth and protoperithecial development, as well as a reduced percentage of cell death during HI. These observations suggest that altering these five sites results in a VIB-1 protein that is not fully functional (hypomorphic allele). Our genetic and phenotypic data show a genetic interaction between *ime-2* and *vib-1* during protoperithecial development and during HI and protease secretion in response to nitrogen starvation and for the localization and processing of a mitochondrial matrix protein. These observations suggest a complex regulatory interaction between these two proteins in a number of cellular functions. Further experiments will unravel the interconnection of these two proteins and the signaling and transcriptional regulatory pathways they regulate in different cellular contexts.

Acknowledgments

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Literature Cited

- Aanen, D. K., A. J. Debets, N. L. Glass, and S. J. Saupe, 2010 Biology and genetics of vegetative incompability in fungi, pp. 274–288 in *Cellular and Molecular Biology of Filamentous Fungi*, edited by K. A. Borkovich, and D. J. Ebbole. ASM Press, Washington, DC.
- Abadjieva, A., P. Hilven, K. Pauwels, and M. Crabeel, 2000 The yeast *ARG7* gene product is autoproteolyzed to two subunit peptides, yielding active ornithine acetyltransferase. J. Biol. Chem. 275: 11361–11367.
- Abdelwahid, E., S. Rolland, X. Teng, B. Conradt, J. M. Hardwick et al., 2011 Mitochondrial involvement in cell death of nonmammalian eukaryotes. Biochim. Biophys. Acta 1813: 597–607.
- Abe, H., and C. Shimoda, 2000 Autoregulated expression of *Schizosaccharomyces pombe* meiosis-specific transcription factor Mei4 and a genome-wide search for its target genes. Genetics 154: 1497–1508.
- Ahmed, N. T., D. Bungard, M. E. Shin, M. Moore, and E. Winter, 2009 The Ime2 protein kinase enhances the disassociation of the Sum1 repressor from middle meiotic promoters. Mol. Cell. Biol. 29: 4352–4362.
- Amoutzias, G. D., Y. He, J. Gordon, D. Mossialos, S. G. Oliver *et al.*, 2010 Posttranslational regulation impacts the fate of duplicated genes. Proc. Natl. Acad. Sci. USA 107: 2967–2971.
- Benjamin, K. R., C. Zhang, K. M. Shokat, and I. Herskowitz, 2003 Control of landmark events in meiosis by the CDK Cdc28 and the meiosis-specific kinase Ime2. Genes Dev. 17: 1524–1539.
- Bhattacharyya, R. P., A. Remenyi, B. J. Yeh, and W. A. Lim, 2006 Domains, motifs, and scaffolds: the role of modular interactions in the evolution and wiring of cell signaling circuits. Annu. Rev. Biochem. 75: 655–680.
- Borkovich, K. A., L. A. Alex, O. Yarden, M. Freitag, G. E. Turner et al., 2004 Lessons from the genome sequence of *Neurospora* crassa: tracing the path from genomic blueprint to multicellular organism. Microbiol. Mol. Biol. Rev. 68: 1–108.
- Bowman, B. J., M. Draskovic, M. Freitag, and E. J. Bowman, 2009 Structure and distribution of organelles and cellular location of calcium transporters in *Neurospora crassa*. Eukaryot. Cell 8: 1845–1855.
- Brush, G. S., N. A. Najor, A. A. Dombkowski, D. Cukovic, and K. E. Sawarynski, 2012 Yeast *IME2* functions early in meiosis upstream of cell cycle-regulated SBF and MBF targets. PLoS ONE 7: e31575.
- Choi, G. H., A. L. Dawe, A. Churbanov, M. L. Smith, M. G. Milgroom et al., 2012 Molecular characterization of vegetative incompatibility genes that restrict hypovirus transmission in the chestnut blight fungus *Cryphonectria parasitica*. Genetics 190: 113–127.
- Chu, S., and I. Herskowitz, 1998 Gametogenesis in yeast is regulated by a transcriptional cascade dependent on Ndt80. Mol. Cell 1: 685–696.

- Chu, S., J. DeRisi, M. Eisen, J. Mulholland, D. Botstein *et al.*, 1998 The transcriptional program of sporulation in budding yeast. Science 282: 699–705.
- Colot, H. V., G. Park, G. E. Turner, C. Ringelberg, C. M. Crew et al., 2006 A high-throughput gene knockout procedure for *Neurospora* reveals functions for multiple transcription factors. Proc. Natl. Acad. Sci. USA 103: 10352–10357.
- Crabeel, M., A. Abadjieva, P. Hilven, J. Desimpelaere, and O. Soetens, 1997 Characterization of the *Saccharomyces cerevisiae ARG7* gene encoding ornithine acetyltransferase, an enzyme also endowed with acetylglutamate synthase activity. Eur. J. Biochem. 250: 232–241.
- Cybis, J., and R. H. Davis, 1975 Organization and control in the arginine biosynthetic pathway of *Neurospora*. J. Bacteriol. 123: 196–202.
- Degterev, A., and J. Yuan, 2008 Expansion and evolution of cell death programmes. Nat. Rev. Mol. Cell Biol. 9: 378–390.
- Dementhon, K., G. Iyer, and N. L. Glass, 2006 VIB-1 is required for expression of genes necessary for programmed cell death in *Neurospora crassa*. Eukaryot. Cell 5: 2161–2173.
- Deshmukh, K., K. Anamika, and N. Srinivasan, 2010 Evolution of domain combinations in protein kinases and its implications for functional diversity. Prog. Biophys. Mol. Biol. 102: 1–15.
- Espagne, E., P. Balhadere, M. L. Penin, C. Barreau, and B. Turcq, 2002 HET-E and HET-D belong to a new subfamily of WD40 proteins involved in vegetative incompatibility specificity in the fungus *Podospora anserina*. Genetics 161: 71–81.
- Fleissner, A., A. R. Simonin, and N. L. Glass, 2008 Cell fusion in the filamentous fungus, *Neurospora crassa*. Methods Mol. Biol. 475: 21–38.
- Glass, N. L., and K. Dementhon, 2006 Non-self recognition and programmed cell death in filamentous fungi. Curr. Opin. Microbiol. 9: 553–558.
- Glass, N. L., and I. Kaneko, 2003 Fatal attraction: nonself recognition and heterokaryon incompatibility in filamentous fungi. Eukaryot. Cell 2: 1–8.
- Glass, N. L., J. Grotelueschen, and R. L. Metzenberg, 1990 Neurospora crassa A mating-type region. Proc. Natl. Acad. Sci. USA 87: 4912–4916.
- Hall, C., J. Welch, D. J. Kowbel, and N. L. Glass, 2010 Evolution and diversity of a fungal self/nonself recognition locus. PLoS ONE 5: e14055.
- Hickey, P. C., S. R. Swift, M. G. Roca, and N. D. Read, 2004 Livecell imaging of filamentous fungi using vital flourescent dyes and confocal microscopy. Methods Microbiol. 34: 63–87.
- Hirsh, H. M., 1954 Environmental factors influencing the differentiation of protoperithecia and their relation to tyrosinase and melanin formation in *Neurospora crassa*. Physiol. Plant. 7: 72–92.
- Holt, L. J., J. E. Hutti, L. C. Cantley, and D. O. Morgan, 2007 Evolution of Ime2 phosphorylation sites on Cdk1 substrates provides a mechanism to limit the effects of the phosphatase Cdc14 in meiosis. Mol. Cell 25: 689–702.
- Honigberg, S. M., 2004 Ime2p and Cdc28p: co-pilots driving meiotic development. J. Cell. Biochem. 92: 1025–1033.
- Honigberg, S. M., and K. Purnapatre, 2003 Signal pathway integration in the switch from the mitotic cell cycle to meiosis in yeast. J. Cell Sci. 116: 2137–2147.
- Hutchison, E., S. Brown, C. Tian, and N. L. Glass, 2009 Transcriptional profiling and functional analysis of heterokaryon incompatibility in *Neurospora crassa* reveals that reactive oxygen species, but not metacaspases, are associated with programmed cell death. Microbiology 155: 3957–3970.
- Hutchison, E. A., and N. L. Glass, 2010 Meiotic regulators Ndt80 and Ime2 have different roles in *Saccharomyces* and *Neurospora*. Genetics 185: 1271–1282.

- Irniger, S., 2011 The Ime2 protein kinase family in fungi: more duties than just meiosis. Mol. Microbiol. 80: 1–13.
- Kaneko, I., K. Dementhon, Q. Xiang, and N. L. Glass, 2006 Nonallelic interactions between *het-c* and a polymorphic locus, *pin-c*, are essential for nonself recognition and programmed cell death in *Neurospora crassa*. Genetics 172: 1545–1555.
- Kassir, Y., N. Adir, E. Boger-Nadjar, N. G. Raviv, I. Rubin-Bejerano et al., 2003 Transcriptional regulation of meiosis in budding yeast. Int. Rev. Cytol. 224: 111–171.
- Katz, M. E., K. A. Gray, and B. F. Cheetham, 2006 The *Aspergillus nidulans xprG* (*phoG*) gene encodes a putative transcriptional activator involved in the response to nutrient limitation. Fungal Genet. Biol. 43: 190–199.
- Keeping, A., D. Deabreu, M. Dibernardo, and R. A. Collins, 2010 Gel-based mass spectrometric and computational approaches to the mitochondrial proteome of *Neurospora*. Fungal Genet. Biol. 48: 526–536.
- Kosti, I., Y. Mandel-Gutfreund, F. Glaser, and B. A. Horwitz, 2010 Comparative analysis of fungal protein kinases and associated domains. BMC Genomics 11: 133.
- Kothe, G. O., and S. J. Free, 1998 The isolation and characterization of *nrc-1* and *nrc-2*, two genes encoding protein kinases that control growth and development in *Neurospora crassa*. Genetics 149: 117–130.
- Lafontaine, D. L., and M. L. Smith, 2012 Diverse interactions mediate asymmetric incompatibility by the *het-6* supergene complex in *Neurospora crassa*. Fungal Genet. Biol. 49: 65–73.
- Macho, A., D. Decaudin, M. Castedo, T. Hirsh, S. A. Susin *et al.*, 1996 Chloromethyl-X-Rosamine is an aldehyde-fixable potential-sensitive fluorochrome for the detection of early apoptosis. Cytometry 25: 333–340.
- Madeo, F., E. Herker, S. Wissing, H. Jungwirth, T. Eisenberg et al., 2004 Apoptosis in yeast. Curr. Opin. Microbiol. 7: 655–660.
- Madeo, F., D. Carmona-Gutierrez, J. Ring, S. Buettner, T. Eisenberg *et al.*, 2009 Caspase-dependent and caspase-independent cell death pathways in yeast. Biochem. Biophys. Res. Commun. 382: 227–231.
- Maheshwari, R., and A. Navaraj, 2008 Senescence in fungi: the view from *Neurospora*. FEMS Microbiol. Lett. 280: 135–143.
- Margolin, B. S., M. Freitag, and E. U. Selker, 1997 Improved plasmids for gene targeting at the *his-3* locus of *Neurospora crassa* by electroporation. Fungal Genet. Newsl. 44: 34–36.
- McCluskey, K., 2003 The Fungal Genetics Stock Center: from molds to molecules. Adv. Appl. Microbiol. 52: 246–262.
- Micali, C. O., and M. L. Smith, 2006 A nonself recognition gene complex in *Neurospora crassa*. Genetics 173: 1991–2004.
- Moore, M., M. E. Shin, A. Bruning, K. Schindler, A. Vershon et al., 2007 Arg-Pro-X-Ser/Thr is a consensus phosphoacceptor sequence for the meiosis-specific Ime2 protein kinase in Saccharomyces cerevisiae. Biochemistry 46: 271–278.
- Moses, A. M., and C. R. Landry, 2010 Moving from transcriptional to phospho-evolution: generalizing regulatory evolution? Trends Genet. 26: 462–467.
- Newmeyer, D., 1970 A suppressor of the heterokaryon-incompatibility associated with mating type in *Neurospora crassa*. Can. J. Genet. Cytol. 12: 914–926.
- Obenauer, J. C., L. C. Cantley, and M. B. Yaffe, 2003 Scansite 2.0: proteome-wide prediction of cell signaling interactions using short sequence motifs. Nucleic Acids Res. 31: 3635–3641.
- Pak, J., and J. Segall, 2002a Regulation of the premiddle and middle phases of expression of the *NDT80* gene during sporulation of *Saccharomyces cerevisiae*. Mol. Cell. Biol. 22: 6417– 6429.
- Pak, J., and J. Segall, 2002b Role of Ndt80, Sum1, and Swe1 as targets of the meiotic recombination checkpoint that control exit from pachytene and spore formation in *Saccharomyces cerevisiae*. Mol. Cell. Biol. 22: 6430–6440.

- Paoletti, M., and C. Clave, 2007 The fungus-specific HET domain mediates programmed cell death in *Podospora anserina*. Eukaryot. Cell 6: 2001–2008.
- Perkins, D. D., 1988 Main features of vegetative incompatibility in *Neurospora crassa*. Fungal Genet. Newsl. 35: 44–46.
- Pittenger, T. H., 1957 The mating type alleles and heterokaryon formation in *Neurospora crassa*. Microbiol. Genet. Bull. 15: 21–22.
- Poot, M., Y. Z. Zhang, J. A. Kramer, K. S. Wells, L. J. Jones *et al.*, 1996 Analysis of mitochondrial morphology and function with novel fixable fluorescent stains. J. Histochem. Cytochem. 44: 1363–1372.
- Read, N. D., A. Fleissner, M. G. Roca, and N. L. Glass, 2010 Hyphal fusion, pp. 260–273 in *Cellular and Molecular Biology of Filamentous Fungi*, edited by K. A. Borkovich, and D. J. Ebbole. ASM Press, Washington, DC.
- Radu, M., and J. Chernoff, 2009 The DeMSTification of mammalian Ste20 kinases. Curr. Biol. 19: R421–R425.
- Ruepp, A., A. Zollner, D. Maier, K. Albermann, J. Hani *et al.*, 2004 The FunCat, a functional annotation scheme for systematic classification of proteins from whole genomes. Nucleic Acids Res. 32: 5539–5545.
- Sarkar, S., G. Iyer, J. Wu, and N. L. Glass, 2002 Nonself recognition is mediated by HET-C heterocomplex formation during vegetative incompatibility. EMBO J. 21: 4841–4850.
- Schmidhauser, T. J., F. R. Lauter, V. E. Russo, and C. Yanofsky, 1990 Cloning, sequence, and photoregulation of *al-1*, a carotenoid biosynthetic gene of *Neurospora crassa*. Mol. Cell. Biol. 10: 5064–5070.
- Sharon, A., A. Finkelstein, N. Shlezinger, and I. Hatam, 2009 Fungal apoptosis: function, genes and gene function. FEMS Microbiol. Rev. 33: 833–854.
- Shin, M. E., A. Skokotas, and E. Winter, 2010 The Cdk1 and Ime2 protein kinases trigger exit from meiotic prophase in *Saccharo-myces cerevisiae* by inhibiting the Sum1 transcriptional repressor. Mol. Cell. Biol. 30: 2996–3003.
- Shiu, P. K., and N. L. Glass, 1999 Molecular characterization of tol, a mediator of mating-type-associated vegetative incompatibility in *Neurospora crassa*. Genetics 151: 545–555.
- Shubassi, G., N. Luca, J. Pak, and J. Segall, 2003 Activity of phosphoforms and truncated versions of Ndt80, a checkpoint-regulated sporulation-specific transcription factor of *Saccharomyces cerevisiae*. Mol. Genet. Genomics 270: 324–336.
- Smith, H. E., and A. P. Mitchell, 1989 A transcriptional cascade governs entry into meiosis in *Saccharomyces cerevisiae*. Mol. Cell. Biol. 9: 2142–2152.
- Sopko, R., S. Raithatha, and D. Stuart, 2002 Phosphorylation and maximal activity of *Saccharomyces cerevisiae* meiosis-specific transcription factor Ndt80 is dependent on Ime2. Mol. Cell. Biol. 22: 7024–7040.
- Strudwick, N., M. Brown, V. M. Parmar, and M. Schroder, 2010 Ime1 and Ime2 are required for pseudohyphal growth of *Saccharomyces cerevisiae* on nonfermentable carbon sources. Mol. Cell. Biol. 30: 5514–5530.
- Tait, S. W., and D. R. Green, 2010 Mitochondria and cell death: outer membrane permeabilization and beyond. Nat. Rev. Mol. Cell Biol. 11: 621–632.
- Tian, C., T. Kasuga, M. S. Sachs, and N. L. Glass, 2007 Transcriptional profiling of cross pathway control in *Neurospora crassa* and comparative analysis of the Gcn4 and CPC1 regulons. Eukaryot. Cell 6: 1018–1029.
- Townsend, J. P., and D. L. Hartl, 2002 Bayesian analysis of gene expression levels: statistical quantification of relative mRNA level across multiple strains or treatments. Genome Biol. 3: RESEARCH0071.
- Vogel, H. J., 1956 A convenient growth medium for *Neurospora*. Microbiol. Genet. Bull. 13: 42–46.

- Westergaard, M., and H. K. Mitchell, 1947 A synthetic medium favoring sexual reproduction. Am. J. Bot. 34: 573–577.
- Winter, E., 2012 The Sum1/Ndt80 transcriptional switch and commitment to meiosis in *Saccharomyces cerevisiae*. Microbiol. Mol. Biol. Rev. 76: 1–15.
- Xiang, Q., and N. L. Glass, 2002 Identification of *vib-1*, a locus involved in vegetative incompatibility mediated by *het-c* in *Neurospora crassa*. Genetics 162: 89–101.
- Xiang, Q., and N. L. Glass, 2004 The control of mating type heterokaryon incompatibility by *vib-1*, a locus involved in *het-c*

heterokaryon incompatibility in Neurospora crassa. Fungal Genet. Biol. 41: 1063.

- Xu, L., M. Ajimura, R. Padmore, C. Klein, and N. Kleckner, 1995 NDT80, a meiosis-specific gene required for exit from pachytene in *Saccharomyces cerevisiae*. Mol. Cell. Biol. 15: 6572–6581.
- Zhang, Z., and J. P. Townsend, 2010 The filamentous fungal gene expression database (FFGED). Fungal Genet. Biol. 47: 199–204.

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Diversification of a Protein Kinase Cascade: IME-2 Is Involved in Nonself Recognition and Programmed Cell Death in *Neurospora crassa*

Elizabeth A. Hutchison, Joanna A. Bueche, and N. Louise Glass

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Figure S1 Phenotype of *vib-1* phosphorylation site mutants. (A) wild type (FGSC 2489), (B) $\Delta vib-1$ (FGSC 11309), (C) transformant with *vib-1*^{S60A}-gfp mutant allele (strain 1XA) and (D) transformant with *vib-1*^{S60D}-gfp mutant allele (strain 1XD). Strains expressing VIB-1^{S60A}-GFP (E) and VIB-1^{S60D}-GFP (F) exhibit normal nuclear localization (shown in panels at right). Brightfield images of hyphae are shown in corresponding panels to the left.



Figure S2 Average growth rate (mm/day) of strains grown on minimal media in race tubes. Asterisks (*) indicate strains with growth rates significantly different (p<0.05) than the WT strain. Strains used are listed in table 1 and are as follows: WT (FGSC 2489), $\Delta vib-1$ (FGSC 11309), $\Delta ime-2$ (FGSC 17937), $\Delta vib-1 \Delta ime-2$ (DVI.4).



Figure S3 Formation of *N. crassa* protoperithecia after 7 days of growth on water agar. Numbers of protoperithecia for each strain are shown as a percentage of wild type. Strains used are listed in table 1 and are as follows: WT (FGSC 2489), $\Delta vib-1$ (FGSC 11309), $\Delta ime-2$ (FGSC 17937), $\Delta vib-1 \Delta ime-2$ (DVI.4).

Table S1 Mitochondrial genes

Gene	Annotation ^a	Decreased expression levels in Δime -
		2
NCU_16001	NADH dehydrogenase subunit 2	
NCU_16002	conserved hypothetical	yes
NCU_16003	cytochrome c oxidase subunit 3	yes
NCU_16004	NADH dehydrogenase subunit 6	
NCU_16005	mitochondrial ribosomal protein S5	yes
NCU_16006	NADH dehydrogenase subunit 2	
NCU_16007	NADH dehydrogenase subunit 3	yes
NCU_16008	NADH dehydrogenase subunit 4L	yes
NCU_16009	hypothetical protein	
NCU_16010	laglidadg endonuclease	
NCU_16011	hypothetical protein	
NCU_16012	NADH dehydrogenase subunit 5	
NCU_16013	cytochrome b	
NCU_16014	group I intron endonuclease	yes
NCU_16015	laglidadg endonuclease	yes
NCU_16016	cytochrome c oxidase subunit 1	yes
NCU_16017	hypothecial protein	yes
NCU_16018	NADH dehydrogenase subunit 1	yes
NCU_16019	group I intron endonuclease	
NCU_16020	NADH dehydrogenase subunit 4	yes
NCU_16021	laglidadg endonuclease	
NCU_16022	hypothetical protein	
NCU_16023	hypothetical protein	
NCU_16024	ATPase subunit 8	yes
NCU_16025	ATPase subunit 6	
NCU_16026	laglidadg endonuclease	
NCU_16027	ATPase subunit 9	yes
NCU_16028	cytochrome c oxidase subunit 2	

^aBroad Institute

NCU #	Annotation
NCU00101	PBN-1
NCU00258	40S ribosomal protein S7
NCU00269	Histone-lysine N-methyltransferase, H3 lysine-36 specific
NCU00276	DNA polymerase gamma, mitochondrial
NCU00311	Protein VTS1
NCU00552	Phytoene dehydrogenase; albino 1 protein
NCU00675	Protein EFR-3
NCU01157	Glutamate-cysteine ligase
NCU01206	Histone-lysine N-methyltransferase, H3 lysine-4 specific
NCU01312	Myb-like DNA-binding protein MYB-1
NCU01797	Serine/threonine-protein kinase NRC-2 (Non-repressible conidiation protein 2)
NCU01954	Pre-mRNA-splicing factor CWC-24
NCU02466	SVP1-like protein 2
NCU03678	Protein SSH-4
NCU03725	Transcription factor VIB-1
NCU03819	COPII coat assembly protein SEC-16
NCU03894	Serine/threonine-protein kinase STE-20
NCU04190	Probable exocyst complex component SEC-8
NCU04191	Protein RAI-1
NCU05589	Protein SIP-5
NCU05876	pH-response regulator protein PAL-A/RIM-20
NCU06276	Increased rDNA silencing protein 4
NCU06498	Pre-mRNA-splicing factor CWC-21
NCU07693	Protein STU-1
NCU07863	Chromatin modification-related protein VID-21
NCU08566	Nuclear distribution protein nudE homolog 1 (RO-11)
NCU08699	Putative oxidoreductase BLI-4, mitochondrial
NCU09068	Nitrogen catabolic enzyme regulatory protein
NCU09730	Kinesin heavy chain
NCU09993	Helicase SWR-1

 Table S2
 Potential N. crassa IME-2 targets for phosphorylation, predicted by Scansite^{a, b}

^a <u>http://scansite.mit.edu/;</u> ^b one mismatch was allowed at the final consensus position

GENETICS

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NCU_16003	cytochrome c oxidase subunit 3	yes
NCU_16004	NADH dehydrogenase subunit 6	
NCU_16005	mitochondrial ribosomal protein S5	yes
NCU_16006	NADH dehydrogenase subunit 2	
NCU_16007	NADH dehydrogenase subunit 3	yes
NCU_16008	NADH dehydrogenase subunit 4L	yes
NCU_16009	hypothetical protein	
NCU_16010	laglidadg endonuclease	
NCU_16011	hypothetical protein	
NCU_16012	NADH dehydrogenase subunit 5	
NCU_16013	cytochrome b	
NCU_16014	group I intron endonuclease	yes
NCU_16015	laglidadg endonuclease	yes
NCU_16016	cytochrome c oxidase subunit 1	yes
NCU_16017	hypothecial protein	yes
NCU_16018	NADH dehydrogenase subunit 1	yes
NCU_16019	group I intron endonuclease	
NCU_16020	NADH dehydrogenase subunit 4	yes
NCU_16021	laglidadg endonuclease	
NCU_16022	hypothetical protein	
NCU_16023	hypothetical protein	
NCU_16024	ATPase subunit 8	yes
NCU_16025	ATPase subunit 6	
NCU_16026	laglidadg endonuclease	
NCU_16027	ATPase subunit 9	yes
NCU_16028	cytochrome c oxidase subunit 2	

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NCU #	Annotation
NCU00101	PBN-1
NCU00258	40S ribosomal protein S7
NCU00269	Histone-lysine N-methyltransferase, H3 lysine-36 specific
NCU00276	DNA polymerase gamma, mitochondrial
NCU00311	Protein VTS1
NCU00552	Phytoene dehydrogenase; albino 1 protein
NCU00675	Protein EFR-3
NCU01157	Glutamate-cysteine ligase
NCU01206	Histone-lysine N-methyltransferase, H3 lysine-4 specific
NCU01312	Myb-like DNA-binding protein MYB-1
NCU01797	Serine/threonine-protein kinase NRC-2 (Non-repressible conidiation protein 2)
NCU01954	Pre-mRNA-splicing factor CWC-24
NCU02466	SVP1-like protein 2
NCU03678	Protein SSH-4
NCU03725	Transcription factor VIB-1
NCU03819	COPII coat assembly protein SEC-16
NCU03894	Serine/threonine-protein kinase STE-20
NCU04190	Probable exocyst complex component SEC-8
NCU04191	Protein RAI-1
NCU05589	Protein SIP-5
NCU05876	pH-response regulator protein PAL-A/RIM-20
NCU06276	Increased rDNA silencing protein 4
NCU06498	Pre-mRNA-splicing factor CWC-21
NCU07693	Protein STU-1
NCU07863	Chromatin modification-related protein VID-21
NCU08566	Nuclear distribution protein nudE homolog 1 (RO-11)
NCU08699	Putative oxidoreductase BLI-4, mitochondrial
NCU09068	Nitrogen catabolic enzyme regulatory protein
NCU09730	Kinesin heavy chain
NCU09993	Helicase SWR-1

 Table S2
 Potential N. crassa IME-2 targets for phosphorylation, predicted by Scansite^{a, b}

^a <u>http://scansite.mit.edu/;</u> ^b one mismatch was allowed at the final consensus position