Control of development, secondary metabolism and light-dependent carotenoid biosynthesis by the velvet complex of *Neurospora crassa*

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Abstract

*Neurospora crassa* is an established reference organism to investigate carotene biosynthesis and light regulation. However, there is little evidence of its capacity to produce secondary metabolites. Here, we report the role of the fungal-specific regulatory velvet complexes in development and secondary metabolism (SM) in *N. crassa*. Three velvet proteins VE-1, VE-2, VOS-1 and a putative methyltransferase LAE-1 show light-independent nucleocytoplasmic localization. Two distinct velvet complexes, a heterotrimeric VE-1/VE-2/LAE-1 and a heterodimeric VE-2/VOS-1 are found in vivo. The heterotrimeric-complex, which positively regulates sexual development and represses asexual sporulation, suppresses siderophore coprogen production under iron starvation conditions. The VE-1/VE-2 heterodimer controls carotene production. VE-1 regulates the expression of more than 15% of the whole genome, comprising mainly regulatory and developmental features. We also studied inter-genera functions of the velvet complex through complementation of *A. nidulans* veA, velB, laeA, vosA mutants with their *N. crassa* orthologues ve-1, ve-2, lae-1 and vos-1, respectively. Expression of VE-1 and VE-2 in *A. nidulans* successfully substitutes the developmental and SM functions of VeA and VelB by forming two functional chimeric velvet complexes in vivo, VelB/VE-1/LaeA and VE-2/VeA/LaeA, respectively. Reciprocally, expression of veA restores the phenotypes of the *N. crassa* ve-1 mutant. All *N. crassa* velvet proteins heterologously expressed in *A. nidulans* are localized to the nuclear fraction independent of light. These data highlight the conservation of the complex formation in *N. crassa* and *A. nidulans*. However, they also underline the inter-genera similarities and differences of velvet roles according to different life styles, niches and ontogenetic processes.
Introduction
Regulation of growth and development in response to environmental stimuli is an essential process in biology. Associated with morphogenetic programmes, cells synchronize growth and metabolism in order to keep up with fluctuating environmental conditions. These stimuli are transduced to the nucleus to alter gene expression by signal transduction complexes composed of receptors, membrane associated protein kinases, mitogen activated protein kinases (MAPK) cascades and downstream regulatory protein complexes (YU AND KELLER 2005; BAYRAM AND BRAUS 2012; FRAWLEY et al. 2018). A central complex in fungi is the heterotrimeric velvet complex that alters gene expression in response to environmental signals such as light, which results in different morphogenetic programs and production of secondary metabolites (SARIKAYA-BAYRAM et al. 2015).

Fungi produce small bioactive compounds also named secondary metabolites (SMs) that have wide-ranging influences on cellular physiology such as antibiotics, mycotoxins, siderophores, antiviral and cytotoxic molecules (KELLER et al. 2005; BRAKHAGE 2013). Each fungus can potentially produce up to 50-100 SMs depending on the genus. The SM genes are often clustered and production of SMs is coordinately controlled by regulatory protein complexes in response to environmental stimuli such as light, carbon source, starvation and pH (KELLER et al. 2005; BRAKHAGE 2013).

Neurospora crassa and Aspergillus nidulans are two model filamentous fungi with different characteristics and life styles, which have been used to understand fundamental questions for eukaryotic molecular genetics (GALAGAN et al. 2003; BORKOVICH et al. 2004; GALAGAN et al. 2005; PARK AND YU 2012; FULLER et al. 2016). However, it has been intriguing whether the regulatory protein complexes controlling developmental programmes or SM production have been structurally and functionally conserved.

N. crassa has been used as a model system to study genetics, biochemistry, enzymology, chromatin biology, cell-cell fusion, light responses and circadian rhythms, and development (SPRINGER 1993; FLEISSNER et al. 2008; ROUNTREE AND SELKER 2010; BAKER et al. 2012; ARAMAYO AND SELKER 2013; HURLEY et al. 2015; DUNLAP AND LOROS 2016). N. crassa can undergo three different sporulation pathways: two different asexual conidiation pathways produce macroconidia and microconidia, while a sexual sporulation pathway leads to formation of meiotic ascospores. Macroconidiation involves the formation of hyphal constrictions at the aerial hyphal tip, initially by a budding process while microconidia, on the contrary, are produced from specialized hyphae in a process that involves the emergence of the microconidial bud and its liberation after breaking the cell wall (SPRINGER 1993). N. crassa uses a heterothallic (self-sterile) system, which requires the fusion of two opposite mating types, a and α (PÖGGELER et al. 2006). In brief, sexual development in N. crassa is initiated by the formation of protoperithecia (semiopen fruiting
body, female organ), followed by the fertilization of the protoperithecia by microconidia from the opposite mating type (male hyphae). Fusion of two opposite mating type nuclei within the perithecia results in the formation of the zygote, which undergoes meiosis to generate sexually formed ascospores. *N. crassa* has a set of light receptors for blue, red and green light but it uses mainly blue light as a signal to adjust cellular activities, sporulation and circadian rhythm, and the WC complex as the main photoreceptor (BAKER et al. 2012; FISCHER et al. 2016). The *N. crassa* genome is relatively poor in SM gene clusters (8-10 clusters) in comparison to Aspergillus (>50 clusters) (KJAERBOLLING et al. 2018) and only a few SMs from *N. crassa* have been identified: the antioxidant histidine-derived ergothioneine, the non-ribosomal peptide coprogen and the polyketide o xoalkylresorcylic acid (ORAS) (HUSCHKA et al. 1985; FUNA et al. 2007; BELLO et al. 2012). Coprogen is a siderophore required for chelating iron ions from the environment, and is historically the oldest metabolite identified from *N. crassa* (HUSCHKA et al. 1985). Microbes use siderophores for the utilization of environmental iron sources. Microbial pathogens sequester iron from high affinity iron-binding molecules such as ferritin, lactoferrin and hemoglobin in the blood of mammals (HAAS 2003; HAAS et al. 2008; HAAS 2014). Furthermore, siderophores have significant potential to be used for treatment of various diseases, for drug delivery, for treatment of heavy metal pollution in the environment, and for the production of functional foods (POCSI et al. 2008).

The velvet family of proteins is restricted to fungi and controls fundamental processes, such as development and SM. The heterotrimeric velvet complex is formed by two velvet transcription factors, VeA and VeIB, and the methyltransferase LaeA, and is a key element in the regulation of light-dependent fungal development and SM production in *A. nidulans* (BAYRAM et al. 2008a). Velvet proteins play essential roles in sporulation, pathogenicity, and SM production in different fungi including human pathogens, endophytic fungi, and plant pathogenic fungi (CALVO 2008; SARIKAYA-BAYRAM et al. 2015). *A. nidulans* vegetative hypha differentiates upon reception of environmental signals. In the light, asexual sporulation (conidiation) is promoted, whereas in the dark the sexual developmental programme (formation of closed fruiting bodies named cleistothecia) is activated (ETXEBESTE et al. 2010; RODRIGUEZ-ROMERO et al. 2010; DYER AND O’GORMAN 2012). In contrast to *N. crassa*, *A. nidulans* has evolved a homothallic (self-fertile) system where the hyphae do not need an opposite mating partner (GALAGAN et al. 2003; GALAGAN et al. 2005; PAOLETTI et al. 2007; DYER AND O’GORMAN 2012). In *A. nidulans* under dark conditions, VeA and VeIB form a heterodimer that enters the nucleus in a process mediated by an α-importin, then the VeA/VeIB heterodimer forms a heterotrimeric velvet complex by interacting with the methyltransferase LaeA, which presumably regulates the transcription of genes for sexual development and SM production including polyketide mycotoxin sterigmatocystin. Although,
LaeA was initially identified as a regulator of secondary metabolism, it also regulates fungal development (Bok and Keller 2004; Sarikaya Bayram et al. 2010). LaeA was shown to be a bona fide methyltransferase since it methylates itself on a methionine residue at position 207 (Patananan et al. 2013). However, other target substrates of LaeA are currently not known. VelB forms an additional heterodimeric complex with the velvet protein VosA, which is required for spore viability and stress tolerance (Ni and Yu 2007; Sarikaya Bayram et al. 2010). The VelB/VosA heterodimer activates the transcription of trehalose biosynthetic genes tpsA and orlA, as well as wetA, a gene essential for spore viability and conidiphore morphology (Park et al. 2012b). The structure of the velvet domain in the VelB-VosA heterodimer is similar to the DNA-binding motif of the rel homology domain of NF-kB transcription factors that activate inflammation pathways in mammals (Ahmed et al. 2013). The role of a fourth velvet protein, VelC, is less clear as the mutant shows slightly increased sexual development (Park et al. 2014). In addition to LaeA, VeA participates in the integration of other signals by interacting with other regulatory proteins. VeA interacts physically and functionally with the putative methyltransferase LlmF, the methyltransferase VipC/VapB heterodimer, the Far1-like DNA binding protein VipA, MAPK MpkB, and the red-light receptor phytochrome FphA (Purschwitz et al. 2008; Bayram et al. 2012; Palmer et al. 2013; Sarikaya-Bayram et al. 2014; Rohrig et al. 2017).

Mechanistic insights into the molecular functions of the velvet complex are mainly limited to Aspergilli. N. crassa and A. nidulans are distantly related with an estimated divergence time of 394 million years (Kumar et al. 2017). N. crassa belongs to Sordariomycetes, which comprise many plant and human pathogenic fungi. In agreement with the phylogenetic distance, there are significant biological differences in response to light, sporulation pathways, and mode of reproduction between N. crassa and A. nidulans. Initial studies of the veA orthologue in N. crassa, ve-I, revealed that it was required for repression of asexual conidiation and the regulation of photocarotenogenesis (Bayram et al. 2008b; Olmedo et al. 2010) (Gil-Sánchez et al. In prep). However, the molecular functions of ve-I and the other velvet complex members in N. crassa remain largely unknown. Here, we characterize the complete set of velvet proteins in N. crassa and show that they form a regulatory velvet complex with a key role in the regulation of development and SM. The functional complementation of some of the velvet proteins between N. crassa and A. nidulans highlights the structural conservation of the velvet complex and its general role in fungal development and the regulation of SM.

Materials and Methods
Strains, media and growth conditions
Strains used in this study are listed in Table S1. General genetic procedures and media used in the handling of N. crassa are available through the Fungal Genetics Stock Center (www.fgsc.net) (Mccluskey 2003). N. crassa strains were mainly cultured in Vogel’s
minimal medium (VMM, 2% sucrose w/v) and *A. nidulans* strains were cultured in glucose (1% w/v) minimal medium (GMM) supplemented with vitamins and trace elements. For iron (Fe) starvation experiments, trace elements were prepared without iron. The flasks were treated with HCl and EDTA as described in detail (Kragl et al. 2007; Schrettl et al. 2008).

For light induction of *N. crassa*, a total of 10⁶ conidia were inoculated on 25 ml Vogel’s liquid medium in 90 mm plates which were incubated in complete darkness at 22 °C for 48 h. Then, the cultures were exposed to light for 30, 60, 120, and 300 min. For carotenoid induction, plates were exposed to light during two min and then incubated at 8 °C for 24 h prior to collection. Light exposure with different intensities was obtained with a quartz halogen lamp installed in a slide projector passed through a filter holder with two heat filters and neutral-density filters, as required to obtain the desired light intensity. A control treatment was always kept in the dark. Mycelia were collected, dried on paper, frozen in liquid nitrogen and stored at –80 °C. Unless otherwise stated, protein quantification was performed with the Bradford assay (Bradford 1976).

**Preparation of nuclear and cytoplasmic protein extracts**
A total of 10⁷ conidia were inoculated in 200 ml of VMM (liquid) and allowed to develop for 48 h at 30 °C under agitation. Three different treatments were prepared: (1) continuous light; (2) continuous darkness; and (3) continuous darkness for 47.5 h followed by a 30 min light exposure. Mycelia were fixed by adding 540 µl of formaldehyde for 15 min and then the reaction was stopped by adding glycine (125 mM). Mycelia were then collected by filtration and frozen in liquid nitrogen. Nuclear fraction extractions were performed as developed by (Baum and Giles 1985) with minor modifications by (Schwerdtfeger and Linden 2000; Froehlich et al. 2002). Further details...
were given in supplemental materials (File S15).

**Antibodies used in this study**

Antibodies were used in the following indicated dilutions: Primary antibodies; monoclonal α-FLAG (1:10000, F3165, Sigma), polyclonal α-histone 3 (H3) (1:5000, Abcam, ab1791), monoclonal α-GFP (1:2000, Santa Cruz, B2, Sc-9996), polyclonal α-human influenza hemagglutinin (HA) (1:4000, Abcam, ab17910), monoclonal α-Tubulin (1:200, Santa Cruz, Sc-32293). Secondary antibodies; HRP conjugated goat α-mouse IgG (1:10000, BioRad, #1721011) was used for α-FLAG and α-Tubulin detections. Goat α-mouse IgG (1:2000, BioRad, #1706516) was used for α-GFP and Goat α-rabbit IgG (1:2000, BioRad, #1706515) was used for α-SkpA, α-HA and α-H3 detections.

**Co-immunoprecipitations of the velvet complexes in N. crassa**

The strains coexpressing VE-1FLAG VE-2HA, VE-1FLAG LAE-1HA, and VE-2FLAG LAE-1HA were grown for 48 h as described for the cellular fractionation experiments and the same three treatments were carried out: continuous light, complete darkness, and complete darkness followed by a 30 min light exposure. Mycelia were also fixed with formaldehyde and collected as described. Mycelial pads (8-10 g wet weight) were harvested by filtration, frozen in liquid nitrogen, and ground to a fine powder which were mixed in a 50 ml tube with 20 ml of IPB buffer [50 mM Tris-HCl pH 7.5, 150 mM NaCl, 1.5 mM MgCl₂, 0.1% (v/v) NP-40] with protease inhibitors (Roche cOmplete™ Protease Inhibitor Cocktail) and were dissolved by vortexing on ice. After mixing, crude extracts were centrifuged at 14,300 × g, at 4°C for 20 min. Supernatants were collected and transferred to ultra-centrifuge tubes. Centrifugation was performed at 169,000 × g at 4°C for 45 min. Then, a 1 ml fraction of each supernatant was transferred to a new tube, labelled “INPUT”, frozen in liquid nitrogen, and stored at –80 °C until use. The remaining supernatant was collected in 15 ml screw-cap tubes and kept at 4°C. During the 45 min centrifugation, 300 µl of M2-FLAG agarose beads (Sigma) were equilibrated by washing them three times with 1 ml of IPB buffer followed by centrifugation at 1,000 × g, 4 °C. After equilibration, 150 µl were added to the supernatant in 15 ml tubes and the mix was incubated for 3 h at 4 °C in a rotating mixer. Beads were then precipitated by centrifuging at 1,500 × g for 10 min at 4 °C. A 1 ml sample of each supernatant were transferred to a new tube, labelled “flowthrough” (FT), frozen in liquid nitrogen, and stored at –80 °C. The beads were washed twice with 10 ml of IPB buffer by gentle rotation for 4 min at 4 °C and finally suspended with 500 µl of IPB buffer and were transferred to new 2 ml tubes. Beads were washed twice with 1 ml IPB buffer by centrifugation at 1,000 × g, 4 °C. At the end, 120 µl of NuPAGE buffer (Invitrogen) was added to the beads, vortexed, and incubated for 10 min at 72 °C with moderate shaking. The supernatant was collected by centrifuging at 1,000 × g, 4 °C and was transferred to new tubes labelled “IP”. Samples were treated with DTT (50 mM) and incubated again for 10 min at 72 °C. Protein concentrations from the INPUT samples were measured in a
Nanodrop instrument (Absorbance set to 280 nm). INPUT, FT and IP samples were used for western blot as described previously. Three independent experiments were performed.

**GFP-TRAP and LC-MS/MS protein identifications**

Immunoprecipitation (IP) of VE-1\(^{\text{GFP}}\), VE-2\(^{\text{GFP}}\), VOS-1\(^{\text{GFP}}\), and LAE-1\(^{\text{GFP}}\) were mainly performed as described in detail (BAYRAM et al. 2012) by using GFP-TRAP magnetic beads (Chromotek). Further details were given in supplemental material (File S15).

**Microscopy**

Spinning disc confocal microscopy of *N. crassa* and *A. nidulans* cells were performed as described earlier (BAYRAM et al. 2012; DETTMANN et al. 2012). Details are given in supplemental material (File S15).

**Secondary metabolite analysis**

**Thin layer chromatography for sterigmatocystin**

Extraction and running of mycotoxin sterigmatocystin (ST) from *A. nidulans* on silica thin layer chromatography plates (Macherey-Nagel) were carried out as described in detail (BAYRAM et al. 2009).

**RP-HPLC analysis**

RP-HPLC analysis was carried out using an Agilent Series 1200 HPLC System with a diode array (DAD) and separation across a water: acetonitrile gradient with 0.1% (v/v) TFA. Gradient conditions of 5-70 % over 30 min at 2 ml/min were used for coprogen analysis with 100 µl injection. Gradient conditions of 5–100% acetonitrile over 30 min at 2 ml/min were used for organic extract analysis with 40 µl injection. Separation was carried out on a C18 column (Agilent Zorbax Eclipse XDB-C18 Semi-Preparative; 5 µm particle size; 9.4 × 250 mm) with DAD detection at 254 and 440 nm. Ferri-coprogen detection was carried out at 440 nm. The peak associated with coprogen was identified in previous analysis by LC-MS/MS detection of a fraction-collected peak. For quantification of coprogen, integration areas detected by HPLC were normalized to biomass (mAUs/ g mycelia).

**LC-MS/MS analysis**

For LC-MS/MS analysis of peaks from RP-HPLC analysis, peaks were collected by fractionation. Fractions were evaporated to dryness in a Speedivac and resuspended in water formic acid (0.1%). Samples were diluted 1 in 50 in water formic acid (0.1%) and spin filtered (0.2 µm) before LC-MS/MS with 1 µl injection. LC-MS/MS analysis was carried out on a nanoflow Agilent 1200 LC system and subjected to tandem mass spectrometry using an Agilent 6340 Ion Trap LC-MS System (Agilent Technologies). Samples were applied to a Zorbax SB-C18 HPLC-Chip with a 40 nl enrichment column and a 75 µm x 43mm (5 µm particle and 300 Å pore size) analytical column. Identification of ferri-coprogen was confirmed via detection of a single charged ion in the MS spectrum (M: 821.3, [M+H]\(^{+}\): observed m/z 822.1; expected m/z 822.3).

**Carotenoid analysis**

Carotenoid analysis was performed as described earlier (CASTRILLO et al. ...
Details were given in supplemental materials (File S15).

**RNA isolation and library preparation**
RNA isolation, poly-A selection and library preparation was done as described in (KLOCKO et al. 2016). The sequencing data were uploaded to the Galaxy web platform, and the public server at usegalaxy.org was used to analyze the data (AFGAN et al. 2016). Specifically, the read quality was checked using FastQC, trimmed the adaptors using Trim Galore, aligned the reads and obtain read count using RNA STAR with intron length set to max 1000 bp. Differential gene expression analysis was done by DESeq2 (LOVE et al. 2014) to examine pair-wise differences between WT and Δve-1 after 48 hour growth under standard or iron-free conditions. Genes with log2 ≥ 1.0 or ≤ −1.0 changed expression and adjusted P values ≤ 0.05 were used to create Venn diagrams using Biovenn (HULSEN et al. 2008). Genes were functionally annotated with Pfam domains and Gene Ontology terms using InterProScan (JONES et al. 2014).

**Statistical Analysis**
Most of the experiments were performed as three independent biological replicates and numerical data are expressed as the mean ± SD and standard error. The mean data were compared for significant differences via the student t-test by using the software Graphpad Prism Version 6.

**Data availability**
Strains and plasmids are available upon request. RNA-seq data was deposited in the Gene Expression Omnibus public database (https://www.ncbi.nlm.nih.gov/geo/) under the accession number GSE123783. Supplemental files are available at FigShare. Table S1 Fungal (Aspergillus nidulans and Neurospora crassa) strains used in this study. Table S2 Oligonucleotides used in this study. Table S3 Plasmids used and constructed in this study. File S1 Interaction partners of VE-1\(^{\text{GFP}}\) fusion in veA\(^{\Delta}\) strain of A. nidulans. File S2 Interaction partners of VE-2 \(^{\text{GFP}}\) fusion in velB\(^{\Delta}\) strain of A. nidulans. File S3 Interaction partners of VOS-1\(^{\text{GFP}}\) fusion in vosI\(^{\Delta}\) strain of A. nidulans. File S4 Interaction partners of LAE-1\(^{\text{GFP}}\) fusion in laeI\(^{\Delta}\) strain of A. nidulans. File S5 Interaction partners of VE-1\(^{\text{GFP}}\) fusion in N. crassa. File S6 Interaction partners of VE-2\(^{\text{GFP}}\) fusion in N. crassa. File S7 Interaction partners of VOS-1\(^{\text{GFP}}\) fusion in N. crassa. File S8 Interaction partners of LAE-1\(^{\text{GFP}}\) fusion in N. crassa. File S9 List of predicted secondary metabolite gene clusters in N. crassa. File S10 List of up- and downregulated genes in ve-1 mutant under standard and iron starvation conditions. File S11 List of GO terms for upregulated genes in ve-1 mutant in both conditions. File S12 List of GO terms for downregulated genes in ve-1 mutant in both conditions. File S13 List of differentially expressed putative secondary metabolite genes in ve-1 mutant in both conditions. File S14 Top twenty up and downregulated genes under iron starvation and standard conditions in ve-1 mutant. File S15 Supplemental materials and methods.

**Results**
N. crassa VE-1 and VE-2 are functional orthologs of A. nidulans VeA and VelB
In addition to the characterized *N. crassa* VE-1 protein (NCU01731; 554 aa, 52% identical to *A. nidulans* VeA (BAYRAM et al. 2008b)), which contains a velvet superfamily domain in its N-terminus with a bipartite nuclear localization signal (NLS), the *N. crassa* genome encodes three additional velvet domain proteins as well as an ortholog of *A. nidulans* LaeA (Figure 1A and (OJEDA-LÓPEZ et al. 2018)). VE-2 (NCU02775; 470 aa; 47% identical to *A. nidulans* VelB) bears a velvet superfamily domain at its C-terminus, and like VelB in *A. nidulans*, lacks an obvious NLS. In contrast, VOS-1 (NCU05964; 447 aa; 44% identical to *A. nidulans* VosA) possesses both an NLS sequence at its C-terminus and a velvet superfamily domain at its N-terminus. The domain structure of VE-3 (NCU07553) is similar to (40% identical) VelC but it appears to have a truncated velvet domain (Figure 1A and (OJEDA-LÓPEZ et al. 2018)). Sequence analysis of LAE-1 (NCU00646; 326 aa; 42% identical to *A. nidulans* LaeA) revealed an S-adenosyl methionine (SAM) binding site and a methyltransferase domain at the centre of the protein.

In order to determine if the four *N. crassa* velvet and the lae-1 genes are functional orthologues of the *A. nidulans* genes, the ve-2, vos-1, and lae-1 genes were heterologously expressed under the control of the constitutive *gpdA* promoter in the *A. nidulans* mutants ΔvelB, ΔvosA and ΔlaeA. In addition, ve-1 was expressed under control of its own native promoter in the *A. nidulans* ΔveA strain (Figure 1). The *N. crassa* ve-1 and ve-2 genes complemented the developmental defects of the *A. nidulans* strains that were characterized by loss of fruiting bodies (cleistothecia), brownish pigment secreted into media and reduced asexual sporulation (Figure 1B). Furthermore, ve-2 complemented the spore inviability (Figure 1C) and the defects of the ΔvelB mutant to oxidative stress (H$_2$O$_2$). However, the *N. crassa* vos-1 and lae-1 genes did not restore the mutant phenotypes of the *A. nidulans* ΔvosA and ΔlaeA strains, i.e. reduced spore viability and small cleistothecia formation defects, respectively. Moreover, we examined if the *N. crassa* velvet genes and lae-1 complemented the defect in the production of the mycotoxin sterigmatocystin (ST) observed in the *A. nidulans* mutants (Figure 1D). Expression of ve-1 and ve-2 in *A. nidulans* restored the lack of ST production in veA and velB mutants to the levels observed in the wild-type, as previously observed for the developmental defects. In contrast, expression of vos-1 and lae-1 did not lead to any significant changes in the chemical phenotypes of the ΔvosA and ΔlaeA strains.

As a further confirmation of the conserved nature of the velvet proteins we complemented the *N. crassa* Δve-1 mutant with the *A. nidulans* veA gene. The two most evident phenotypes of Δve-1 – reduced carotenoid accumulation and reduced growth of aerial hyphae (BAYRAM et al. 2008b) – were restored by the heterologous expression of the *A. nidulans* veA gene fused to the FLAG tag in *N. crassa* (Figure 1E). Furthermore, expression and localization of VeA in *N. crassa* were examined in response to light. The accumulation of VeA did not change under prolonged exposure to light (Figure 1F). VeA protein levels were
mainly increased in the dark and exhibited nuclear localization both under light and dark cultures (only dark is shown) (Figure 1G-H).

In summary our results indicate that the *N. crassa* ve-1 and ve-2 genes are functional orthologues of *veA* and *velB* in *A. nidulans*, while *vos-1* and *lae-1*, seem not to be conserved. This stresses the similarities and differences between these two distantly-related fungal species even in proteins that are known to form complexes in order to function.

*N. crassa* VE-1, VE-2, VOS-1 and LAE-1 accumulate in *A. nidulans* nuclei, but only VE-1 and VE-2 interact with components of the heterotrimeric velvet complex in *A. nidulans* during vegetative growth

In *A. nidulans*, VeA displays light-dependent nuclear localization (STINNETT et al. 2007; BAYRAM et al. 2008a). In order to determine the subcellular localization of *N. crassa* VE-1 and VE-2 in *A. nidulans*, we heterologously expressed the two velvet proteins fused to synthetic green fluorescent proteins (sGFP) under the control of the *gpdA* promoter and examined their localization in cultures of complemented *A. nidulans* mutants grown under light and dark conditions. We did not observe any effect of light on the localization of the fusion proteins, and therefore, only dark grown cultures are shown (Figure 2A, B). *N. crassa* VE-1<sup>GFP</sup> was localized in *A. nidulans* nuclei while the VE-2<sup>GFP</sup> was evenly distributed between nuclei and cytoplasm, resembling the subcellular localization of the corresponding *A. nidulans* proteins (BAYRAM et al. 2008a). Interestingly, VOS-1<sup>GFP</sup> had the strongest nuclear localization whereas LAE-1<sup>GFP</sup> was found both in the nucleus and in the cytoplasm as shown after co-localization with co-labelled red nuclei. Expression of all GFP fusions of VE-1, VE-2, VOS-1 were clearly detected in total protein extracts, confirming that all velvet family proteins and LAE-1 were properly expressed (Figure 2B).

The *A. nidulans* VeA/VelB/LaeA complex controls sexual development and SM production. In addition, VelB forms a heterodimer with VosA that is required for spore viability and trehalose biogenesis. In order to determine if the *N. crassa* velvet proteins and LAE-1 have the capacity to interact with other members of the velvet proteins in *A. nidulans*, VE-1<sup>GFP</sup>, VE-2<sup>GFP</sup>, VOS-1<sup>GFP</sup> and LAE-1<sup>GFP</sup> were immunoprecipitated and interacting proteins were identified using liquid chromatography-mass spectrometry (LC-MS) (Figure 2C-D, File S1 to S4). We detected the presence of VE-1 or VE-2 in chimeric heterotrimeric complexes (VE-1/VelB/LaeA and VeA/VE-2/LaeA) in vegetatively grown cultures of *A. nidulans*. However, we did not detect a chimeric VE-2/VosA complex and none of the velvet complex members were identified after immunoprecipitation of VOS-1<sup>GFP</sup> from vegetative cultures. Consistently, VosA was not identified by LC-MS in the *N. crassa* velvet protein precipitates. We detected the interaction between LAE-1<sup>GFP</sup> and VeA, but not VelB (File S4). In summary, these results suggest that only VE-1 and VE-2 from *N. crassa* form functional chimeric heterotrimeric velvet complexes with their counterparts in *A. nidulans*. 
The *N. crassa* velvet proteins play different roles in asexual and sexual reproduction

In order to understand the molecular and developmental functions of the velvet proteins and the putative methyltransferase LAE-1 in the biology of *N. crassa*, strains with deletions in each of these genes were subjected to development and growth tests (Figure 3). Δve-1, Δve-2, Δvos-1 or Δlae-1 deletion strains did not show any significant changes in growth rate (Figure 3A, B). However, Δve-1 and Δve-2 exhibited reduced aerial hyphae in slants when compared to wild-type or to Δvos-1 and Δlae-1 (Figure 3A). Conidiation increased 2-fold in Δve-1 and Δve-2 compared to wild-type despite their reduced aerial growth, while Δvos-1 and Δlae-1 displayed slight or moderate reductions in conidiation (Figure 3A). The viability of conidia from the mutants did not change significantly. Sexual development was altered in all velvet and the lae-1 mutants, and protoperithecia production was reduced to ~50% of wild-type. Fertilization of the remaining protoperithecia yielded viable ascospores, indicating a functional sexual cycle (Figure 3C-D). In summary, the developmental alterations in the mutants revealed that VE-1 and VE-2 are required for the promotion of aerial hyphal growth and the repression of asexual spore formation. Moreover, VE-1, VE-2 and LAE-1 are important for the production of protoperithecia during sexual development. VOS-1 and LAE-1 play a minor role in the regulation of conidiation.

Velvet proteins control production of secondary metabolites and carotenoid accumulation

The *N. crassa* genome contains ten putative SM gene clusters (File S9), and *N. crassa* produces several SMs including coprogen and ergothioneine (TOTH et al. 2009; BELLO et al. 2012; BELLO et al. 2014). In order to understand the role of the velvet proteins and of LAE-1 on the production of SMs, wild-type and deletion strains were grown in Vogel’s MM medium or in medium lacking iron to promote the production of non-ribosomal peptide (NRPs) siderophores. Iron-free (Fe-free) medium revealed a difference in SM production in the mutants, which was evident by the lighter colors of Δve-1 and Δve-2 liquid cultures (Figure 4A). Growth in Fe-free media resulted in at least 4 to 5-fold elevated levels of coprogen in Δve-1 and Δve-2 in comparison to wild-type, suggesting that ve-1 and ve-2 are negative regulators of coprogen production (Figure 4B). Δlae-1 showed a slight increase in coprogen production, whereas the Δvos-1 mutant did not show any significant changes in coprogen production.

Comparison of organic extracts from 1-week old cultures of wild-type and mutant strains showed at least six unique compounds that significantly changed their accumulation in the mutants when compared to wild-type (Figure 4C; only the Δve-1 chromatogram is shown): A (RT ~18 min), B (RT ~19 min), C (RT ~21 min), D (RT ~23 min), E (RT ~25 min), and F (RT ~32 min). Deletion of ve-1 resulted in a major change in the accumulation of SMs, as it had a significant increase in the abundance of compounds B (p <
0.01), C (p < 0.01), D (p < 0.001), E (p < 0.01), and F (p < 0.01), and a decrease in the abundance of compound A. Deletion of the other genes led to minor changes in the accumulation of a smaller number of SMs. In the Δve-2 mutant, accumulation of compounds C and D increased, while accumulation of compound A decreased; Δlae-1 showed increased accumulation of compounds D and F, and in Δvos-1 decreased accumulation of compound D was observed.

The Δve-1 mutant accumulated less carotenoids than wild-type, and it affected the light regulation of carotenoid biosynthesis (BAYRAM et al. 2008b; Gil-Sánchez et al. in prep.) (Figure 4A). We observed a light-orange color also in Δve-2, suggesting a role of VE-2 in carotenoid biosynthesis. Therefore, we measured accumulation of total carotenoids from all mutants under different light intensities, with the exception of Δve-1 (Figure 4E) (Gil-Sánchez et al. in prep.). The accumulation of carotenoids after light exposure was altered in all mutants, but the effect was more pronounced in Δve-2 and Δlae-1. Both mutants showed a reduction in the accumulation of carotenoids with high intensities of light (>100 J/m²). Our results suggest that VE-1, VE-2 and, partially, LAE-1 are involved in production of several unknown SMs, the siderophore coprogen, and carotenoids.

VE-1, VE-2, VOS-1 and LAE-1 are found in both nuclei and cytoplasm in a light-independent manner
In order to identify the subcellular localization of the velvet proteins and of LAE-1 in N. crassa, GFP-fused versions of each protein were expressed under the control of the ccg-1 promoter at the his-3 locus. VE-1GFP displayed a nucleocytoplasmic localization in both apical and subapical N. crassa cells (Figure 5A). The VE-2GFP fusion was also found homogenously distributed in both nuclear and cytoplasmic fractions in apical and subapical cells. However, VE-2GFP showed a higher abundance in the cytoplasm (Figure 5B, E). LAE-1GFP had a similar nucleocytoplasmic pattern of distribution (Figure 5D, F). However, VOS-1GFP exhibited a more pronounced nuclear localization than the other two velvet proteins (Figure 5C), which is comparable to VosA localization in A. nidulans (NI AND YU 2007). Since VE-2 and LAE-1 exhibited weak nuclear localization, we created strains with fusions of VE-2 or LAE-1 with a triple FLAG tag and expressed the fused genes under their native promoters. The accumulation of the VE-2FLAG or LAE-1FLAG were examined in cytoplasmic and nuclear fractions of cultures grown in dark and light conditions (Figure 5E, F). We detected both proteins in nuclear and cytoplasmic fractions, but VE-2FLAG was less abundant in the nucleus when compared to the LAE-1FLAG. The nuclear subpopulation of LAE-1FLAG showed a higher molecular weight than cytoplasmic LAE-1FLAG, but the nature of the presumptive posttranslational modification was not explored further. The exposure to light did not change the fractionation patterns of VE-2FLAG or LAE-1FLAG, confirming our localization experiments with GFP fusions of these proteins. Thus, the patterns of the velvet proteins in N. crassa showed that they are in both nuclei and cytoplasm, but their localization in the N. crassa nuclei suggests that these proteins might have
important regulatory functions controlling gene expression.

**Trimeric VE-1/VE-2/LAE-1 and dimeric VE-2/VOS-1 complexes co-exist in N. crassa**

Next, we investigated whether the four *N. crassa* velvet proteins and LAE-1 interact to form regulatory protein complexes. Co-immunoprecipitation experiments (CoIPs) of VE-1, VE-2 and LAE-1 were performed using 3XFLAG- and HA-tagged fusion proteins expressed in VMM during vegetative growth under native conditions (Figure 6A). VE-1FLAG co-precipitated both VE-2HA and LAE-1HA. Furthermore, VE-2FLAG also co-precipitated LAE-1HA, consistent with formation of a heterotrimeric velvet complex *in vivo*, as shown in *A. nidulans* (BAYRAM et al. 2008a).

To further support these results, and to identify additional protein interactions of the velvet proteins and of LAE-1, the GFP fusions of the velvet and LAE-1 proteins were used to identify by LC-MS their interacting proteins *in vivo* during vegetative growth (Figure 6B-C). Protein extracts of the vegetative cultures were subjected to GFP-TRAP (pull-down). Silver-stained SDS-PAGE analysis of the GFP-Trap experiment confirmed the molecular sizes of the fusion proteins (Figure 6B). Co-purifying proteins were identified by LC-MS, which suggested that VE-1 interacted with VE-2, but also with VOS-1, presumably via VE-2, which is similar to *A. nidulans* velvet complex (BAYRAM et al. 2008a) (Figure 6C, File S5). Interestingly, VE-2 reciprocally copurified VE-1 and also VOS-1. As a confirmation, VOS-1 reciprocally interacted with VE-2 but not with VE-1, indicating the presence of VE-2/VOS-1 heterodimers. Although LAE-1 was not found in purifications of VE-1 and VE-2, LAE-2GFP purifications led to identification of both VE-1 and VE-2, confirming the CoIP results (Figure 6C, File S5 to S8). In addition, we detected the interaction between the importin DIV-45, a homolog of the yeast karyopherin KAP114p, involved in the nuclear import of specific proteins (MOREHOUSE et al. 1999), and VE-2 and VOS-1.

These results confirmed that the *N. crassa* velvet proteins form at least two distinct protein complexes, a heterotrimeric VE-1/VE-2/LAE-1 complex containing the putative LAE-1 methyltransferase (analogous to VeA/VelB/LaeA in *A. nidulans*), and a heterodimeric VE-2/VOS-1 (analogous to VelB/VosA) complex. The association of VE-2 and VOS-1 with DIV-45 suggest a role for this importin in the transport of the heterodimer and/or the velvet complex into the nucleus.

**VE-1 controls expression of genes required for conidiation, secondary metabolite production and carotenoid biosynthesis**

Our results suggest that VE-1 and VE-2 are the important members of the velvet protein family in *N. crassa*. They form a complex with LAE-1, and corresponding mutants show similar phenotypes in conidiation, sexual fruiting body formation and SM production. In order to identify genes that are under their transcriptional control during different developmental and metabolic processes, we characterized the transcriptome of Δve-1 by RNA-seq experiments. We
cultured wild-type and Δve-1 for 48 h in the presence or absence of iron since the fungus activates SM gene expression in cultures older than 24 h. The genes showing changes in mRNA accumulation at \( \log_2 \geq 1 \) were considered as up and down regulated. A total of 2,927 genes showed changes in mRNA accumulation under either of the two conditions (wild-type vs Δve-1 Vogel’s and iron-free Vogel’s media) (Figure 7, File S10 to S14), which corresponds to approximately 30% of \( N. crassa \) genes.

Under standard growth conditions (with iron), expression of 1,166 genes was downregulated and 1,031 genes was upregulated (21% of genome), whereas under iron-free conditions expression of 907 genes was downregulated and 641 genes was upregulated (15% of genome) in ve-1 compared to wild-type.

When we focused on the set of common genes that were upregulated (226) and downregulated (571) in the presence or absence of iron in Δve-1, we detected enrichment for the following biological processes: oxidation-reduction (11 genes up; 27 genes down), metabolic (9 up), transmembrane (8 up; 18 down), translation (7), transport (4), carbohydrate metabolism (3 up; 17 down), transcription (3 up; 16 down), nucleotide biosynthesis (2 up; 12 down) and other (27 up) were among the upregulated genes under both standard and iron free conditions in Δve-1.

The \( A. nidulans \) genome encodes more than ten LaeA-like methyltransferases (\( llmA \) to \( llmJ \)), which either control development or SM production (PALMER et al. 2013; SARIKAYA-BAYRAM et al. 2014). Two \( N. crassa \) orthologs of the \( A. nidulans \) LaeA-like methyltransferase, encoding NCU10101(\( llmG \)) and NCU10761 (\( llmA \)), were upregulated strongly under iron starvation conditions in Δve-1 (Table 1). In addition to these two methyltransferases, two sugar transporters, a trehalase enzyme, cytochrome c and an endonuclease were among the top ten highly expressed genes (see also the top twenty list in File S14). Under standard growth conditions, mainly oxidation-reduction enzymes (luciferase-like monooxygenase) were at the top of the upregulated list along with major facilitator superfamily genes (Table 2).

Oxidation-reduction, proteolysis, bacterial cell wall degrading enzyme domain encoding (LysM domain) genes were the most drastically downregulated genes under iron starvation conditions in Δve-1 mutant (Table 3). Interestingly, vos-1 was highly, and ve-2 was moderately, downregulated under normal conditions in the Δve-1 mutant, suggesting that VE-1 is required for proper expression of vos-1 and ve-2 (Table 4). In addition to the \( llmG \) and \( llmA \) homologues, which were upregulated in the Δve-1 mutant, the genes for many other methyltransferases, including NCU05841(\( llmB \) (vipC)), two \( llmB \)-like methyltransferase NCU05501 and NCU05832, an O-methyltransferase domain encoding gene NCU05855, two Lae-like genes NCU00304 and NCU00451, were downregulated under both standard and iron starvation conditions (File S14).
The \(\Delta ve-1\) mutant shows increased conidiation and reduced carotenoid production. In agreement with these phenotypes, conidiation genes \(con-6\), and \(con-10\) were moderately upregulated even in submerged media, which precludes conidiation of \(N. crassa\). The carotenoid biosynthetic gene \(al-2\) (phytoene synthase) as well as \(cao-1\) (carotenoid oxygenase) were moderately downregulated under standard conditions in the \(ve-1\) mutant. Expression of the \(N. crassa\) clock components (\(wc-1\), \(wc-2\), \(frq\), \(vvd\)) was unaltered in the absence of VE-1 (File S10).

There are 10 predicted SM gene clusters with 94 genes in the \(N. crassa\) genome. 36 out of 94 genes showed changes in the accumulation of the corresponding mRNAs (either up or downregulation) in either of two conditions (Figure 8A). Expression of SM genes were mainly downregulated in \(\Delta ve-1\) mutant irrespective of iron starvation. Clusters 2, 9 and 10 were mostly downregulated whereas clusters 4 and 8 were upregulated. Expression of the backbone enzymes of cluster 4 (PKS-like), 6 (NRPS), 7 (PKS) were particularly increased. Cluster 6 is responsible for production of coprogen with backbone enzyme NCU07119. Two genes belonging to coprogen production were upregulated in \(\Delta ve-1\) strain after 48 h of growth.

These results reveal that VE-1 regulates more than 20% of the genome under different growth conditions. Furthermore, a great proportion of small methyltransferases, velvet genes, conidiation and carotenoid biosynthetic genes and SM genes require VE-1 for proper expression.

Discussion

Eukaryotes have developed different strategies to coordinate development and cellular physiology in response to environmental conditions. The velvet complex is one of the regulatory protein complexes that fungi use for the coordination of growth, development and SM production. In this study, we have examined the cross-genus functions of the velvet complex by comparing its function in two model fungi from two different genera at the biochemical, cellular and genetical levels.

The two \(N. crassa\) velvet genes \(ve-1\) and \(ve-2\) successfully complemented the deficiencies of the \(A. nidulans\) \(veA\) and \(velB\) mutants, indicating that VE-1 and VE-2 are functional orthologues of VeA and VelB. Intriguingly, VE-1 and VE-2 are functionally incorporated and form heterotrimeric chimeric complexes with either \(A. nidulans\) VelB/LaeA or VeA/LaeA, respectively. However, neither \(N. crassa\) VOS-1 or LAE-1 could complement the functions of \(vosA\) and \(laeA\) in \(A. nidulans\). This might be due to the abolished/reduced ability of VOS-1 and LAE-1 to integrate into the \(A. nidulans\) complex. A residual interaction of \(N. crassa\) LAE-1 was observed with \(A. nidulans\) VeA but not VelB, while we did not observe complexes formed with VOS-1 in the vegetative growth stage. We did not test for interaction partners of VOS-1 in \(A. nidulans\) during asexual sporulation. Therefore, it is still possible that \(N. crassa\) VOS-1 might recruit \(A. nidulans\) VelB during asexual development. Nevertheless, \(N. crassa\) VOS-1 and LAE-1 were expressed and localized
correctly within the *A. nidulans* nuclei. It is intriguing that *N. crassa* LAE-1 could interact with VeA in *A. nidulans*, but did not complement the functions of LaeA, presumably because the LAE-1/VeA heterodimer was not able to recruit VelB. In *A. nidulans*, VeA acts as a bridge between VelB and LaeA. Furthermore, in the absence of LaeA, VeA recruits a high level of VelB (*SARIKAYA BAYRAM et al. 2010*). Presumably the interaction of LAE-1 with VeA interferes with the VeA/VelB interaction in *A. nidulans*, consistent with the possibilities that LAE-1 binds and masks the domain on VeA where VelB binds or changes the conformation of VeA, preventing VelB binding.

Our results allowed us to compare the current model for the mechanism of action of the velvet complexes in *A. nidulans* and *N. crassa* (Figure 8B). The *A. nidulans* velvet complex is formed in the nucleus after the entry of VeA/VelB heterodimers via the importin KapA. In *N. crassa*, however, another importin, DIV-45, which is an orthologue of the yeast karyopherine KAP114p, copurified with both VE-2 and VOS-1 and thus may promote nuclear translocation of the velvet proteins. Since VE-2 interacts with VE-1, this heterodimer is likely to use the DIV-45 importin for nuclear import as well. In *A. nidulans*, the VelB/VeA heterodimer recruits the LaeA methyltransferase upon nuclear entry, and the final heterotrimeric velvet complex participates in the regulation of sexual development, ST production and hülle cell formation to protect growing fruiting bodies. VelB, in addition, forms a heterodimer with VosA that is essential for final spore maturation, spore viability, and trehalose accumulation.

In *N. crassa*, VE-1, VE-2, and LAE-1 form the heterotrimeric velvet complex. However, the VE-1/VE-2 heterodimer has more prominent roles in development than LAE-1, which is only required for full production of protoperithecia. Both VE-1 and VE-2 are equally important for the repression of conidiation and for the activation of carotenoid biosynthesis by light. This phenotypic observation was further confirmed by our transcriptomic results that showed the downregulation of two carotenoid biosynthetic genes. VE-2 forms a heterodimer with VOS-1, which presumably represses asexual conidiation. The slightly reduced conidiation in the Δvos-1 mutant and the minor alterations in SM and carotenoid accumulation suggest a minor role for VOS-1 in the regulation or activities of the velvet complexes.

Few cross-species functional studies with the velvet complex proteins have been reported. In the human pathogenic fungus *A. fumigatus*, ectopic introduction of the *A. nidulans* VeA-TAP fusion resulted in formation of the functional AnVeA-AfVelB-AfLaeA velvet complex (*PARK et al. 2012a*). *veA* is required for production of gliotoxin (GT) and nitrogen source dependent sporulation in *A. fumigatus* (*KRAPPMANN et al. 2005; DHINGRA et al. 2012*). In an inter-genera complementation experiment, the VeA ortholog of the dimorphic fungus *Histoplasma capsulatum*, Vea1, was able to restore cleistothecia formation and partially complemented ST production in *A. nidulans* (*LASKOWSKI-PEAK et al. 2012*). Silencing of VEA1 facilitated the dimorphic switch from
the yeast phase to the mycelial phase in *H. capsulatum*, and virulence of this strain is attenuated in murine and macrophage models.

The VosA/VelB heterodimer of *A. nidulans* has a DNA binding domain that is similar to that of the Rel family of regulatory proteins in the mammalian immune system, and binds to DNA to regulate the transcription of several genes, including sporulation and trehalose genes (Ahmed et al. 2013). VeA was shown to bind to its own promoter (Rauscher et al. 2016). It is likely that VeA acts as a DNA binding domain protein since the velvet domain seems to be important for DNA binding (Sarikaya-Bayram et al. 2015). The *H. capsulatum* velvet family proteins Ryp2 (VelB) and Ryp3 (VosA) are important for the temperature-dependent dimorphic switch and also bind to a large set of *H. capsulatum* genes important for pathogenicity (Beyhan et al. 2013). Not only in *H. capsulatum*, but also in *A. fumigatus*, the veA and laeA members of the velvet complex control temperature-dependent processes like SM production (Lind et al. 2016), underlining the velvet complex uses not only light but also temperature to regulate developmental and metabolic processes in at least two pathogenic fungi.

It is highly likely that the heterodimeric VE-1/VE-2 complex of *N. crassa* binds DNA given the drastic changes in gene expression that we observed after deleting ve-I since the velvet domains were shown to bind to DNA in several other fungi (Ahmed et al. 2013; Beyhan et al. 2013; Becker et al. 2016; Rauscher et al. 2016). A 21% change in the transcriptome was observed under standard growth conditions and 15% under iron starvation conditions when the Δve-I and the wild-type strains were compared (Figure 7). In contrast, loss of *A. fumigatus* veA changes expression of 10% of the genome including more than half of the genes in SM gene clusters (Lind et al. 2016). Interestingly, more than one third of the SM gene clusters whose expression is differentially regulated (14 out of 37) are downregulated in the absence of veA in *A. fumigatus*. In the plant pathogenic fungus, *A. flavus*, expression of 5% of all genes were differentially expressed in a veA mutant (Cary et al. 2015). 28 out of 56 gene clusters contain at least one gene that is either up- or downregulated in veAΔ strain in *A. flavus*. Similarly, in *N. crassa*, expression of almost one third of SM genes (36 out of 94) is either up- or down-regulated in either normal or iron starvation media. Coprogen production increases three-fold when ve-I is missing. However, this metabolic change was not reflected in gene expression, because only two genes belonging to the coprogen cluster (cluster 6) were upregulated. This discrepancy between gene expression and coprogen production may be due to the different time points used for the assay of SM accumulation (7 days) and RNA extraction (48 h) since it is extremely difficult to extract intact RNA from 7 days old cultures in comparison to SMs, which can be easily extracted from old cultures.

In summary, our results suggest a strong conservation in the activities and interactions of the velvet family in the fungal kingdom given the biological activity of chimeric versions of the velvet complexes in *A. nidulans*. In *N.*
crassa, however, our results show how the velvet complex has further evolved and specialized compared to the A. nidulans counterpart since the VE-1/VE-2 heterodimer plays a major regulatory role when compared to the trimeric VE-1/VE-2/LAE-1 complex. The key role of VE-1 in N. crassa biology, as shown by the developmental and metabolic phenotypes of the Δve-1 mutant, further indicates specialization when compared with the phenotype of the veA mutant in A. nidulans and other Aspergilli, and suggests that in-depth understanding of the role of the velvet complex in fungal biology will require detailed characterization in selected model fungi like N. crassa. It will be interesting to know whether or where the velvet proteins, VE-1, VE-2 and VOS-1 bind to in N. crassa genome. Molecular activities of A. nidulans LaeA are currently unknown except for automethylation activity. It will be also intriguing to learn whether N. crassa LAE-1 has similar or different biochemical activities than A. nidulans LaeA.

Fungi are prominent in human activities as they are sources of chemicals used as pharmaceuticals and drugs and include pathogens of plants and animal, including humans. The key role of the velvet proteins in fungal development and SM production suggests that a full understanding of their complex dynamics in fungal biology and metabolism will improve our capacity to use fungi for food and drug production and may mitigate their detrimental effects on humans and plant hosts.

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Figure 1 Interspecies functionality of velvet proteins in Aspergillus nidulans and Neurospora crassa for light-dependent development and secondary metabolite production. (A) Four velvet family proteins are shown encoded by N. crassa genome. Top scale indicates their length in amino acids (aa). VE-1 and VOS-1 contains a nuclear localization signal (NLS). (B) Growth and developmental responses of A. nidulans velvet complex mutants, veAΔ, velBΔ, laeAΔ,
vosAΔ and their complementation by N. crassa velvet orthologs (ve-1, ve-2, vos-1, lae-1) expressed under the control of the native (ve-1) promoter or the constitutive gpdA promoter (ve-2, vos-1, lae-1) in comparison to wild-type (WT) grown under light and dark conditions. 5x10^3 spores were point-inoculated on GMM plates and grown at 37 °C for 5 days. (C) Spore viability and stress tolerance test veAΔ, velBΔ, laeAΔ, vosAΔ strains and for strains expressing corresponding heterologous proteins. Spore viability tests were performed with 10^3 spores at 37 °C. (+) Represents complementations. Asterisks denote the significant difference (P<0.05). (D) Examination of mycotoxin production in the velvet complex mutants and their respective complemented strains by thin layer chromatography (TLC). Strains were grown on GMM plates with oatmeal for 3 days in the dark. Quantification of the ST from two independent biological replicates of TLC plates. STs: Sterigmatocystin standard. (E) Complementation of Δve-1 phenotypes (reduced carotenogenesis, stunted hyphae) by A. nidulans VeA3XFLAG fusion expressed under N. crassa ve-1 promoter. (F) Expression of VeA3XFLAG in N. crassa under constant illumination. 30 µg total protein was loaded per lane. (G) Presence of VeA3XFLAG in total crude extract, cytoplasmic and nuclear fractions (50 µg in each lane) in the light and dark. Strain was grown in liquid VMM for 48 h at 30 °C and then exposed to white light for 30 min or kept in the dark as a control. (H) Cellular localization of VeA GFP expressed under ceg-1 promoter in N. crassa grown in VMM on a coverslip overnight at 30 °C. DAPI stains DNA in the nuclei blue and arrows indicate the nuclear localization of VeA GFP.
**Figure 2** Localization and complex formation of *Neurospora crassa* velvet proteins in *Aspergillus nidulans*. (A) Subcellular localization of VE-1, VE-2, LAE-1 and VOS-1\(^GFP\) fusion proteins in the respective *A. nidulans* velvet mutants (*veA\(\Delta\), *velB\(\Delta\), *laeA\(\Delta\), *vosA\(\Delta\)). Chimeric fungi (10\(^3\) spores) expressing the fusion proteins were grown at 30 °C for 16 h and observed by confocal microscopy. DRAQ5 stains the nuclei red. (B) Protein expression of VE-1\(^GFP\), VE-2\(^GFP\), LAE-1\(^GFP\), VOS-1\(^GFP\) fusion proteins assayed by a western blot analysis. In general, 80 µg total protein of crude cell extracts were loaded per lane but for the highly expressed VE-2 and LAE-1 samples, 40 µg were loaded. α-GFP detects GFP fusions and α-SkpA detects constitutively expressed SkpA protein, which serves as a loading control (therefore less SkpA is seen in the VE-2 and LAE-1 lanes. (C) Silver stained (10%) SDS gels of the GFP TRAP pulldowns of fusion proteins. Arrows indicate the positions of strongly staining full length precipitated fusion proteins. (D) Interaction partners of VE-1, VE-2, VOS-1 and LAE-1 in *A. nidulans* during vegetative growth. The results of GFP TRAP pulldowns digested with trypsin and identified by LC-MS. Gene ID (locus number), name and number of unique peptides found are given in the tables (See Files S1-S4). Chimeric velvet complexes found in each purification are given at the top of the table.
Figure 3 Developmental functions of the velvet and Lae-1 orthologs of *N. crassa* (A) Slant growth phenotypes and hyphae formation of the wild-type along with Δve-1, Δve-2, Δvos-1 and Δlae-1 strains on VMM after 4 days at RT (left panel) and on plates (right panel) under continuous illumination. (B) Quantification of relative growth rates, asexual conidiation and resistance to UV light. Growth rate was measured on VMM at RT for 24 h. Conidiation was measured from plates grown for 4 days. For the UV test, 50 fresh spores spread on sorbose medium plates were exposed to UV for 0, 2, 4, or 8 min, incubated on plates at RT for 2-3 days, and colony forming units were counted. Asterisks denote the significant difference (P<0.05) (C-D) Formation of protoperithecia, perithecia and ascospores of the WT and velvet mutants along with lae-1 mutant. Equal amounts of fungal spores were grown at the center of corn meal agar (CMA) plates for 7 days for protoperithecia formation and 14 days at RT for perithecia maturation and ascospore analysis. Perithecia were squeezed open for photo-microscopy of ascospores.
Figure 4 Regulatory functions of the velvet orthologs in production of coprogen, carotenes and various other compounds. (A) A comparison of WT, ve-1, ve-2, vos-3, and lae-1 mutants of *N. crassa* grown in liquid VMM (left panel) with iron (Fe) and without (iron starvation) (right panel) at 37 °C for 7 days. (B) Production of the iron chelating and growth promoting compound coprogen was normalized to biomass (p < 0.05). (C) Mirror image comparison of the WT and ve-1 mutant’s organic extraction chromatogram by RP-HPLC (254 nm). (D) Production of five compounds significantly changed mainly in ve-1 or ve-2 mutants. Chromatograms were compared with WT and any compounds of interest were quantified by integrating areas associated with each peak. Quantifications were obtained from the supernatants of the three biological replicates grown in liquid iron-starvation Vogel’s media (FVMM) at 37 °C for 7 days. (E) Quantification of total carotene produced by ve-2, vos-1 and lae-1 mutants in comparison to WT under a gradient of illumination.
Figure 5 Expression and subcellular localization of *N. crassa* velvet orthologs. (A) Subcellular localizations of VE-1\(^\text{GFP}\), (B) VE-2\(^\text{GFP}\), (C) LAE-1\(^\text{GFP}\) and (D) VOS-1\(^\text{GFP}\) fusion proteins expressed under the control of the clock-controlled gene-1 (ccg-1) promoter at the his-3 locus of *N. crassa*. White arrows indicate the position of nuclei with GFP signals. Plasma membranes and septa were stained red by a membrane staining FM4-64 dye and nuclei were visualized using Histone 1 (H1)-RFP fusion. (E-F) Nuclear enrichment of VE-2 and LAE-1 triple FLAG fusion proteins expressed under the control of their native promoters extracted from cultures grown under light and dark conditions. Cultures were either grown in the dark 48 h and then exposed to light for 30 min, or they were grown under continuous white light for 48 h. Total, Cyto and Nuc represent total, cytoplasmic and nuclear extracts, respectively. Each lane was loaded with 70 µg protein. α-Tubulin and α-Histone 3 (H3) were used as loading controls for cytoplasmic and nuclear proteins, respectively.
Figure 6 in vivo interactions of the velvet protein complexes in *N. crassa* during vegetative growth (A) Co-immunoprecipitations (CoIPs) of the velvet complex using triple (3X) FLAG and HA-tagged versions of VE-1, VE-2 and LAE-1 expressed under their native promoters during vegetative growth. Triple HA-tagged VE-2 and LAE-1 were precipitated with VE-13XFLAG fusion. HA-tagged LAE-1 was also precipitated with VE-23XFLAG fusion. (B) Silver-stained SDS gels (10%) of the GFP-TRAP pulldowns of VE-1, VE-2, VOS-1 and LAE-1GFP fusion proteins from *N. crassa* grown for 24 hours at 30 °C in liquid Vogel’s MM medium. Arrows indicate the positions of precipitated full-length fusion proteins stained by Silver reagent. (C) Interaction partners of VE-1, VE-2, VOS-1 and LAE-1 after GFP-TRAP-MS in *N. crassa* during vegetative growth. Gene IDs are given as locus number (NCU). NUP; the number of unique peptides identified in MS. LC-MS identifications of interaction partners of respective proteins (See Files S5-S8).
Figure 7 Control of gene expression by ve-l gene. (A) Upregulated genes and biological processes in the ve-l mutant grown under standard VMM and iron-free conditions during 48 h submerged culture. The Venn diagram represents the 805 upregulated genes (red) in the ve-l mutant under standard conditions (Iron containing) and 415 upregulated genes (green) in the ve-l mutant specific to iron-starvation. The overlapping 226 genes represent the genes upregulated under both conditions. Bar charts show the number of GO biological process terms upregulated (226 genes) in both conditions (see Files S10-S14). (B) Downregulated genes and biological processes in the ve-l mutant under standard and iron-free conditions during the 48 h submerged culture. 595 and 336 genes were downregulated in the ve-l mutant in standard and iron starvation conditions, respectively. Bar charts show the number of GO biological process terms downregulated (571 genes) in both conditions.
Figure 8 Expression of secondary metabolite genes and comparative model of the velvet complexes in *A. nidulans* and *N. crassa*. (A) Expression of eight putative secondary metabolite gene clusters of *Neurospora* in liquid 48 hour cultures under standard or iron starvation conditions in an Δve-1 strain in comparison to WT. PKS: Polyketide synthase, NRPS: Nonribosomal peptide synthase. SM cluster numbers, and the “backbone” genes (with NCU numbers indicated) in the clusters are identified. Backbone genes were indicated with an asterisk. Expression of some backbone genes did not change (those without the asterisks) (see S13 Table). (B) Velvet complex has been mostly studied in *A. nidulans* (upper panel). The VeIB-VeA heterodimer, formed by the two velvet family proteins (represented by red and orange spherical shapes), enters into the nucleus with help of importin alpha (KapA, Blue). In the nucleus VeIB-VeA form a heterodimer with the methyltransferase LaeA. This complex has several functions; the VeA-VeB part of the complex is mainly responsible for sexual development and SM production (shown as green lines). In addition, LaeA has a particular function on formation of protective hülle cells required for protection and nursing of growing
fruit bodies. In addition to its role in the velvet complex, VelB is also a part of heterodimeric complex formed with VosA, which activates spore viability, trehalose accumulation (green lines) and also represses asexual conidiation (red lines). In *N. crassa*, the situation is similar. The VE-1/VE-2 or VE-2/VOS-1 heterodimer enters into nucleus, presumably with help of importin DIV-45 (KAP114p ortholog) since DIV-45 was found in purifications of both VE-2 and VOS-1. The VE-2/VE-1 heterodimer is a major player to repress asexual conidiation and SM production including coprogen (red line) and to activate carotene biosynthesis (green line). The VE-1/VE-2/LAE-1 complex is required for maximum protoperithecia formation (sexual development). The VE-2/VOS-1 heterodimer presumably represses conidiation and hyphal growth.
Table 1 Top ten upregulated genes under iron starvation in ve-I mutant

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Table 2 Top ten upregulated genes under standard conditions in ve-1 mutant

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Table 3 Top ten downregulated genes under iron starvation conditions in ve-I mutant

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Table 4 Top ten downregulated genes under standard conditions in ve-1 mutant

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