Evidence of a Demethylase-Independent Role for the H3K4-Specific Histone Demethylases in Aspergillus nidulans and Fusarium graminearum Secondary Metabolism

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Fungi produce a plethora of secondary metabolites (SMs) involved in cellular protection, defense, and signaling. Like other metabolic processes, transcription of SM biosynthesis genes is tightly regulated to prevent an unnecessary use of resources. Genes involved in SM biosynthesis are usually physically linked, arranged in secondary metabolite gene clusters (SMGCs). Research over the last decades has shown that chromatin structure and posttranslational modifications (PTMs) of histones represent important layers of SMGC regulation. For instance, trimethylation of histone H3 lysine 4 (H3K4me3) is a PTM typically associated with promoter regions of actively transcribed genes. Previously, we have shown that the H3K4me3-specific, JmjC domain-containing histone demethylase KdmB functions not only in repression but also in activation of secondary metabolism in Aspergillus nidulans, suggesting that KdmB has additional functions apart from histone demethylation. In this study, we identified demethylase-independent functions of KdmB in transcriptional regulation of SM gene clusters. Furthermore, we show that this activating and demethylase-independent role of the H3K4 demethylase is also conserved in the phytopathogenic fungus Fusarium graminearum. Lack of FgKdm5 resulted in significant downregulation of five of seven analyzed SMs, whereby only one SMGC depends on a functional JmjC-domain. In A. nidulans strains deficient in H3K4 methylation, i.e., cclAΔ, largely phenocopied kdmBΔ, while this is not the case for most of the SMs analyzed in Fusarium spp. Notably, KdmB could not rescue the demethylase function in Δfgkdm5 but restored all demethylase-independent phenotypes.

Keywords: KDM5, histone demethylase, secondary metabolism, Aspergillus nidulans, Fusarium graminearum, H3K4 methylation, CclA, SppA
INTRODUCTION

Filamentous fungi produce a plethora of structurally diverse low molecular weight compounds, so-called secondary metabolites (SMs), that may provide a competitive advantage for the producing organism (Macheleidt et al., 2016). Among SMs are potent toxins that frequently occur in contaminated food and feed (Streit et al., 2013), but also pharmaceuticals and plant hormones that are applied in medicine and agriculture, respectively (Dreyfuss et al., 1976; Anke et al., 1977; Rademacher, 1997; Manzoni and Rollini, 2002). SMs are synthesized by stepwise enzymatic reactions. Enzymes involved in the biosynthesis are usually encoded in physically linked SM gene clusters (SMGCs). To prevent an unnecessary use of resources, the expression of SM genes is tightly regulated. Each SMGC requires defined environmental or developmental conditions to be induced (Macheleidt et al., 2016). One advantage of SM gene clustering could be the possibility to co-regulate them by large-scale chromatin structure that would grant or deny access to underlying regulatory elements (Gacek and Strauss, 2012; Pfannenstiel and Keller, 2019). Chromatin is composed of histones and chromatin-associated factors. Histones are modified by posttranslational chemical modifications (e.g., acetylation, methylation, or phosphorylation) on different residues, and a certain combination of these histone marks generates a recognition platform for chromatin regulators that strongly influence how the underlying genetic information is read (Jenuwein, 2002; Wood et al., 2005; Drelon et al., 2018). The combination of histone PTMs defines the overall chromatin structure and transcriptional activity at a given genomic region. Chromatin-based gene silencing relies on heterochromatin structures that are characterized by densely packed nucleosomes, whereas loosely arranged nucleosomes in euchromatin are open for transcriptional activation. Both chromatin states are associated with the activity of certain histone-modifying enzymes that are often part of large complexes involved in the recognition, addition, or removal of histone PTMs (Kouzarides, 2007; Piunti and Shilatifard, 2016). Notably, many histone PTMs recruit additional chromatin-modifying enzymes that further remodel the chromatin landscape, thereby enabling cells to respond dynamically to environmental or developmental cues (Gacek and Strauss, 2012; Pfannenstiel and Keller, 2019).

One prominent euchromatic histone PTM is trimethylation of lysine 4 on histone H3 (H3K4me3). H3K4 methylation is established by the COMPASS complex (complex of proteins associated with Set1) that is conserved in euarkocytes (Miller et al., 2001; Krogan et al., 2002; Hughes et al., 2004; Lee et al., 2007; Wu et al., 2008; Jiang et al., 2011; Shilatifard, 2012). H3K4 methylation is also associated with active transcription of the majority of genes in Aspergillus spp. and Fusarium spp. (Connolly et al., 2013; Wiemann et al., 2013; Gacek-Matthews et al., 2016). Notably, while H3K4 methylation is generally associated with transcriptional gene activation, COMPASS-mediated silencing of genes located near telomeres has also been described (Briggs et al., 2001; Krogan et al., 2002; Bok et al., 2009; Mikheyeva et al., 2014). In agreement with these findings, deletion of the COMPASS components ccdA and CCL1 (homologous to bre2) in Aspergillus spp. and Fusarium spp., respectively, resulted in increased expression of several SM cluster genes and the subsequent production of additional SMs in these fungi (Bok et al., 2009; Palmer et al., 2013; Shinohara et al., 2016; Studt et al., 2017). Noteworthy, in Aspergillus nidulans as well as in Fusarium graminearum and Fusarium fujikuroi, SMGCs are largely devoid of this histone PTM, even under activated conditions (Connolly et al., 2013; Wiemann et al., 2013; Gacek-Matthews et al., 2016; Studt et al., 2017). H3K4 methylation is countered by enzymes of the KDM5-family of histone demethylases, which are able to remove one or two methyl groups from H3K4me3. Their function has thus mainly been associated with transcriptional repression. However, there is also evidence for transcriptional activation by KDM5-family members, e.g., in mammalian systems and Drosophila melanogaster (Klose et al., 2007; Secombe et al., 2007; Secombe and Eisenman, 2007; Seward et al., 2007). Recently, we have shown that the A. nidulans KDM5 homolog KdmB balances H3K4 trimethylation levels and both positively and negatively influences around one third of the transcriptome under the tested growth conditions. Notably, around 50% of SM-related genes are affected by kdmB deletion, and the vast majority (80%) of them show reduced expression in the kdmB deletion mutant (Gacek-Matthews et al., 2016). This bipartite, locus-dependent activating or repressing function seems to be conserved also in F. fujikuroi (Janevska et al., 2018). Thus, it is intriguing to hypothesize that the activating functions of KdmB are also conserved also conserved in fungi and that SM-related genes are major targets of this multi-domain chromatin regulator.

KDM5 proteins are composed of multiple domains, which are necessary for their diverse role in transcriptional regulation. In detail, KdmB is a member of the Jumonji C (JmjC) domain-containing demethylases, catalyzing lysine demethylation of histones through an oxidative reaction that requires iron (Fe²⁺) and α-ketoglutarate as substrates (Klose et al., 2006). Additionally to its catalytic JmjC-domain (IPR003347), KdmB harbors a putative JmjN domain (IPR003349), an AT-rich interaction domain (ARID, IPR001606), a C5HC2-type zinc finger motif (IPR004198), and two plant homeo domains (PHD, IPR001606). Recently Liu and Secombe (2015) have shown that the KDM5 homolog Lid in D. melanogaster activates gene expression independently of its demethylase domain by recognizing the chromatin context through its PHD domains. In fact, the N-terminal PHD domain is able to read unmethylated lysine residues, whereas the C-terminal PHD domain binds to H3K4me2/me3 marks (Li et al., 2010). Here, we provide evidence for a demethylase-independent role of KdmB in secondary metabolism in A. nidulans. Notably, lack of H3K4 methylation, which was engineered by removal of crucial COMPASS components (CcdA, SetA) or by replacement of H3K4 by arginine (H3K4R), largely phenocopied the kdmB deletion with regard to SMGC regulation and SM production. Furthermore, we show that both the demethylase-dependent and independent role of KDM5-family members in SM gene regulation extends to other fungi like the plant pathogen F. graminearum. However, functional differences must exist between both homologs, as loss of the demethylase function in F. graminearum Δfgkdm5 strains is not rescued by expression of A. nidulans ΔkdmB.
MATERIALS AND METHODS

Fungal Strains, Media, and Growth Conditions

The wild-type strains of A. nidulans (AnWT) pabaA1 veA1 (Pontecorvo et al., 1953) and F. graminearum (FgWT) PH-1 (FGSC 9075, NRRL 31084) were used as parental strains for deletion and targeted mutagenesis experiments. For protoplasting, DNA isolation, and western blot analyses, all strains were grown in darkness for 3 days on solid complete media (CM) (Pontecorvo et al., 1953) covered with cellophane sheets (Folia Bringmann) at 37 and 20°C in case of A. nidulans and F. graminearum, respectively. Fungal growth test were performed either on solid minimal medium (FMM: F. graminearum; AMM: A. nidulans; Todd et al., 2007), complete medium (FCM: F. graminearum; ACM: A. nidulans), or complete medium supplemented with 1 M sorbitol (ACMS: A. nidulans; Govindaraghavan et al., 2014). For A. nidulans gene expression and SM analyses, the respective strains were grown in liquid Aspergillus minimal media (AMM; Todd et al., 2007) in darkness at 37°C for 48 h. In case of F. graminearum, SM analyses were performed with strains grown on potato dextrose agar (PDA) in darkness at 25°C for 2 weeks (Giese et al., 2013). Cultivation of Saccharomyces cerevisiae and Escherichia coli was performed as previously described (Schumacher, 2012).

Plasmid Constructions

If not mentioned otherwise, plasmid generations were performed using yeast recombinational cloning as described by Schumacher (2012). All primers used for polymerase chain reaction (PCR) were obtained from Sigma-Aldrich GmbH. For all generated constructs, the strategy and primers used in this study can be found in Supplementary Tables S1, S2. Generally, for deletion constructs, the 5′ and 3′ flank fragments were amplified with the primer pairs 5F//5R and 3F//3R from genomic DNA of the respective parental wild-type strain (AnWT, FgWT). For A. nidulans constructs, auxothrophic marker cassettes (pyrG or riboB) were used as selectable markers and amplified from A. fumigatus genomic DNA. For F. graminearum the hph or neoR resistance cassette was amplified from template pCSN44 (Staben et al., 2017) and pfgccl-IL (Studt et al., 2017), respectively. Constructs for targeted mutagenesis were generated by using primers harboring the desired mutation. Complementation constructs were generated by amplifying the wild-type gene kdmB/FgKDM5 from parental genomic DNA. In case of A. nidulans, kdmB was driven by its native promoter and terminator sequences, whereas for F. graminearum, the FgKDM5 gene was driven by the native promoter and fused to the glucanase terminator of Botrytis cinerea (BcTgluc) from pNAH-PoliC::bcltf3-gfp (Brandhoff et al., 2017). For cross-complementation constructs, the kdmB gene was amplified from A. nidulans genomic DNA driven by the native Fusarium promoter. The kdmB gene was fused to BcTgluc, followed by neoR amplified from pfgccl-IL (Studt et al., 2017). S. cerevisiae FY834 was transformed with the obtained fragments and the EcoRI/Xhol-restricted pRS426 yielding the plasmids pKdmB*, pKdmBΔ/kdmB*, pCClΔ, pSppΔ, pΔgkdm5, pFgKDM5*, pΔgkdm5/FgKDM5*, and pΔgkdm5/kdmB*α. A proof-reading polymerase was used for amplification of all plasmid parts, and correct assembly of the gained plasmids was verified by restriction digest and/or sequencing. All primers used for plasmid generation and strain generation strategies are listed in Supplementary Tables S1, S2.

Fungal Transformations

In case of A. nidulans, experimental strains were obtained by transformation into an nkuAΔ strain, which reduces the frequency of non-homologous integration (Nayak et al., 2006). Fungal transformations were performed essentially as described (Tilburn et al., 1983; Studt et al., 2017). In case of A. nidulans, the constructs pKdmB* and pKdmBΔ/kdmB*α were linearized with MluI, whereas for pCClΔ and pSppΔ, the knockout fragments were amplified prior transformation with the primers An_CclA_del_1F//An_CclA_del_02R and An_SppA_del_F1//An_SppA_del_R2. In case of F. graminearum, we used the split marker approach (Goswami, 2012). For deletion/mutation of kdm5/FgKDM5 the fragments were amplified from pΔgkdm5/pΔgKdm5* with the primers Kdm5_5F//Split-mark-hphF and Split-mark-hphR/Kdm5_3R. For complementation of Δgkdm5, the plasmid Δgkdm5/FgKDM5 was integrated ectopically into the Δgkdm5 strain. For cross-complementation, the fragments were amplified from pΔgkdm5/kdmB*α with the primers Kdm5_5F//GenIsplitF and GenIsplitR//Kdm5_3R. Transformed protoplasts were regenerated as described by Studt et al. (2017).

In case of A. nidulans KdmB* mutants, homologous recombination events were checked by using the primers ribo_fum_check_F//8211_downst_R (Supplementary Figure S6). For further verification of the mutation H642G and E644Q, a fragment was amplified with the primers F9//R8 and sequenced. For kdmBΔ/kdmB*α mutants, homologous recombination events were verified by using the primers dia_kdmB_F02//kdmB_comp_R02 and dia_pyrG_R01//dia_kdmB_F01 (Supplementary Figure S6). The knock-out mutants cclAΔ and sppAΔ were checked using the primers dia_dCclA_UP_F//dia_pyrG_fum_5 and dia_SppA_UP_F//dia_pyrG_fum_5′ for the upstream region and dia_dCclA_DownR//dia_pyrG_fum_3′ and dia_dSppA_DownR//dia_pyrG_fum_3′ for the downstream region. The absence of the wild-type genes, cclA and sppA, was checked using the primer pairs dia_WT_CclA_F//dia_WT_CclA_R and dia_WT_SppA_F//dia_WT_SppA_R, respectively (Supplementary Figure S2).

In case of F. graminearum, homologous recombination events of the Δgkdm5 mutants were verified by the primer pairs dia_kdm5_F//pCSN44_hph-trpcT as well as dia_kdm5_F//pCSN44_trpc_P2 (Supplementary Figure S8). FgKdm5* mutants were checked with the primer pairs dia_kdm5_F//Kdm5_DM_UP_R for the upstream fragment and dia_Kdm5_R//pKS_GenR_F for the downstream fragment (Supplementary Figure S8). Δgkdm5/kdmB*α mutants were checked with the primer pairs dia_Kdm5_F//KdmB_dia_check_UP_R and dia_Kdm5_R//pks_genR_F (Supplementary Figure S10). Δgkdm5/FgKDM5*α mutants were verified with the primer pair Kdm5_HK_F//Kdm5_HK_R (Supplementary Figure S8).
Standard Molecular Techniques

For DNA isolation, lyophilized mycelium was ground to a fine powder, re-suspended in extraction buffer, and isolated as previously described (Tilburn et al., 1983; Cesin, 1992). Isolated genomic DNA was used for PCR amplification and Southern blot analysis. Deletion, complementation, and cross-complementation fragments were amplified with the proof-reading Phusion High-Fidelity DNA Polymerase (Thermo Fisher Scientific), and PCR reactions were set up accordingly to the manufacturers’ protocol. For diagnostic PCRs, the GoTaq® Green Master Mix (Promega) was used, and the PCR reactions were set up according to the users’ manual. For Southern, genomic DNA of cclAΔ, sppAΔ and Afgkdm5 was digested with the enzymes Xhol, Eco321 and HindIII, respectively. The digested DNA was separated on a 1% (w/v) agarose gel and transferred onto positively charged nylon membranes (Roche Diagnostics GmbH, Germany) by downward blotting. Probes were labeled with DIG-11-dUTP using the DIG-High Prime DNA Labeling and Detection Starter Kit II from Roche. Primers used for probe generation are listed in Supplementary Table S1.

For expression analyses, RNA was extracted from lyophilized mycelium using the TRIzol Reagent (Thermo Fisher Scientific) according to the manufacturers’ instruction. For cDNA synthesis, 1 μg of total RNA was treated with DNaseI (Thermo Fisher Scientific) and subsequently reversely transcribed using the iScript™ cDNA Synthesis Kit (BioRad). Generated plasmids were extracted and purified from E. coli and S. cerevisiae with the GeneJET™ plasmid miniprep kit (Fermentas GmbH, St. Leon-Rot, Germany). Sequencing of each plasmid was performed using primers listed in Supplementary Table S1. For western blot analyses, mycelium from 2 to 3 days old strains was ground to a fine powder, and proteins were extracted as described (Studt et al., 2016). Roughly, 10 μg (A. nidulans) and 15 μg (F. graminearum) of proteins were used for SDS-PAGE and subsequent western blotting. The membrane was probed with 1:4,000 dilutions of H3 C-Term (Active Motif), H3K4me1 (Active Motif, AM61633), H3K4me2 (Active Motif, AM39141), and H3K4me3 (Active Motif, AM 39159 for F. graminearum, Abcam, ab8580 for A. nidulans) primary antibodies and 1:10,000 dilutions of anti-rabbit (Sigma A0545) HRP conjugated secondary antibody. Chemiluminescence was detected with Clarity™ ECL Western Substrate and ChemDoc™ XRS (Bio-Rad). Densitometric quantification of western blot signals was performed with the ImageJ software and normalized to the histone H3 C-term signal. Subsequently, the signal of the according wild type was set to a value 1; consequently, the presented results are the fold change to the control reaction. Each western blot was performed at least three times for each set of samples.

Chemical Analyses

A. nidulans strains were grown for 48 h in liquid AMM under SM inducing conditions (1% glucose, 10 mM NaNO3), and the supernatant was analyzed. The samples were run on a QTrap 5500 LC–MS/MS System (Applied Biosystems, Foster City, CA, USA) equipped with a TurboIonSpray electrospray ionization (ESI) source and a 1290 Series HPLC System (Agilent, Waldbronn, Germany). Chromatographic separation was done at 25°C using a Gemini C18 150 × 4.6 mm i.d., 5 μm particle size, equipped with a C18 3 × 4 mm i.d. security guard cartridge (Phenomenex, Torrance, CA, USA). The chromatographic method and chromatographic and mass spectrometric parameters are described elsewhere (Malachová et al., 2014).

F. graminearum strains were grown for 2 weeks on PDA plates and subsequently extracted using MeOH/CH2Cl2/EtOAc (1/2/3, v/v), evaporated, and resuspended in methanol/H2O (1/1, v/v) as previously described (Sørensen and Sondergaard, 2014). Extracts from the FgWT and two independent fgkdm5 deletion mutants were initially analyzed by high-resolution mass spectrometry in positive ionization mode as previously described (Wollenberg et al., 2016). Quantitative analyzes of known SMs from F. graminearum were performed on a Thermo Vantage triple stage quadrupole mass spectrometer (Thermo Fisher Scientific, San José, CA, USA) with a heated electrospray ionization probe using the chromatography and MS settings for deoxynivalenol, zearalenone, fusaricin, aurofusarin, fusaristatin A, fusarilin H, and chrysogine as previously described (Sørensen and Sondergaard, 2014; Wollenberg et al., 2017).

Reverse Transcriptase-Quantitative Polymerase Chain Reaction

RT-qPCR was performed with iQ SYBR Green Supermix (Bio-Rad, Munich, Germany) using an iCycler iQ Real-Time PCR System (Bio-Rad). To quantify mRNA levels of SMGCs in A. nidulans, the following primers were used: PEN gene cluster (q_ipnA_F//q_ipnA_R), ST gene cluster (q_aflR_F//q_aflR_R), ORS gene cluster (q_orsA_F//q_orsA_R), and the MDP gene cluster (q_mdpG_F//q_mdpG_R). To quantify the expression level of kdmB, the primers q_kdmB_F//q_kdmB_R were used. Levels of mRNA were related to constitutively expressed reference genes, i.e., AN6838 encoding ß-tubulin (q_TUB_F//q_TUB_R) and AN0290 encoding actin (q_ActA_F//q_ActA_R).

In case of F. graminearum, the following primers were used for FgKDM5 expression (q_Kdm5_F//q_Kdm5_R). Levels of mRNA were related to following reference genes: FSFG_06257 encoding glyceraldehyde 3-phosphate dehydrogenase (GAPDH) (GAPDH_qPCR_fwd//GAPDH_qPCR_rev), FSFG_07335 encoding actin (qPCR_actin_F//qPCR_actin_R), and FSFG_09530 encoding ß-tubulin (cdNA_ß-TUB_F//cdNA_ß-TUB_R). Primer efficiencies in the RT-qPCR were kept between 90 and 110%. Relative expression levels were calculated using the ΔΔCt method (Pfaffl, 2001). Experiments were performed in biological and technical duplicates. Primer sequences are listed in Supplementary Table S1.

RESULTS

Lack of COMPASS Components Largely Phenocopies KdmB With Regard to Secondary Metabolite Gene Regulation

In a previous study, we have shown that KdmB regulates the H3K4 methylation levels and plays a central role in secondary metabolism in A. nidulans (Gacek-Matthews et al., 2016). Here, the majority of SMGCs (11 out of 18 differentially expressed SMGCs) are
less expressed in a strain deleted for kdmB compared to the *A. nidulans* wild-type strain (AnWT) under SM cultivation conditions. However, H3K4me3 is under-represented in chromatin landscapes of SMGCs and, if present, H3K4me3 levels are not increased but decreased at these sites in *kdmBΔ*. This suggests that KdmB promotes SM gene expression either by regulating SM genes independently of H3K4me3 or indirectly by targeting genes of trans-acting factors. To study the relevance of H3K4 methylation for KdmB-mediated regulation, we have now analyzed mutants altered in H3K4 methylation (i.e., H3K4R, setAΔ, cclAΔ, sppAΔ) for their characteristics in SM biosynthesis. Strains deficient in H3K4 methylation, i.e., a strain in which H3K4 has been replaced by arginine (H3K4R) as well as a strain lacking SetA, the catalytic subunit of the COMPASS complex homologous to the histone methyltransferase Set1 are very sick (Govindaraghavan et al., 2014). Also in our hands both strains featured retarded hyphal growth (20% of AnWT) on solid Aspergillus complete medium (ACM) and did not grow at all on solid (Supplementary Figure S1) or in liquid Aspergillus minimal medium (AMM, data not shown). Thus, these strains could not rationally be included in the present study to analyze the impact of H3K4 methylation on SM gene regulation in *A. nidulans*.

Two additional COMPASS components have been described to influence the H3K4 methylation status, namely the subunits Bre2 and Spp1, both required for efficient H3K4 trimethylation in *S. cerevisiae* (Roguè et al., 2001; Dehè et al., 2006; Acquaviva et al., 2013). Based on the phenotype of reverse genetic screens, the *Aspergillus* homologs of Bre2 were designated CclA and found to participate in repression of subtelomeric SMGCs in *A. nidulans* (Bok et al., 2009) and *A. fumigatus* (Palmer et al., 2013). The Spp1 homolog in *Aspergillus*, SppA, has only been characterized so far in *A. oryzae* where it is necessary for full H3K4 trimethylation (Shinohara et al., 2016). We identified the Spp1 homolog in the *A. nidulans* genome database by BlastP analysis as AN2850 (E-value of 2.0e-17). We deleted both cclA and sppA in the same genetic background and at least two independent mutants were verified by diagnostic PCR and Southern blot analysis (Supplementary Figure S2). Western analysis with antibodies directed against different methylation forms of H3K4 showed that lack of CclA leads to a severe reduction in mono-, di-, and trimethylation of this residue in *vivo*, whereas the absence of SppA reduced H3K4 tri- and dimethylation by roughly 50% with a concomitant increase of H3K4 monomethylation by a factor of two (Figure 1A). Contrary to set1Δ and H3K4R, the cclAΔ and sppAΔ strains did not show such extreme growth phenotypes. Deletion of sppA did not detectably alter growth characteristics when compared to AnWT, while cclAΔ mutant featured around 50% reduced radial growth on solid complete and minimal media and around 30% reduced biomass formation in liquid AMM shake cultures (Figure 1B; Supplementary Figure S3). To sum up, the strongest phenotype with regard to radial growth was observed for H3K4R followed by setAΔ and cclAΔ, all being significantly impaired in H3K4 methylation, while deletion of sppA had only mild impact on H3K4 methylation levels and growth.

To analyze if strains with reduced or abolished H3K4me3 levels affect secondary metabolism in *A. nidulans* in a similar manner as strains lacking the H3K4 demethylase KdmB, the generated mutant strains were grown under SM-inducing conditions (liquid AMM with 10 mM sodium nitrate) and analyzed for SMs known to be regulated by this chromatin mark. Biosynthesis of both penicillin G and sterigmatocystin were almost abolished in strains lacking CclA. This is similar although not identical to what has been found in *kdmBΔ*. In agreement with the reduced production of the metabolites, also the expression of the biosynthetic genes, i.e., *ipnA* (encoding the non-ribosomal peptide synthetase (NRPS) required for penicillin (PEN) production) and *afIR* (encoding the pathway-specific transcription factor for sterigmatocystin (ST) biosynthesis) were reduced (Figure 1C). This indicates that H3K4 methylation and KdmB are required for PEN and ST biosynthesis. Contrary to cclAΔ, the lack of SppA did not reduce the production of the two model SMs. Notably, the levels of penicillin G (200% of AnWT) and sterigmatocystin (300% of AnWT) were even found to be increased in sppAΔ compared to AnWT (Figure 1C). However, transcript levels for the tested cluster genes, i.e., *ipnA* (PEN cluster) and *afIR* (ST cluster) were basically unchanged in comparison to AnWT indicating that the altered COMPASS complex without the SppA subunit and the concomitantly slightly reduced overall H3K4 methylation levels are not crucial for PEN and ST cluster transcription (Figure 1C).

We also tested to which extent the slightly or severely altered global H3K4 methylation levels in cclAΔ and sppAΔ mutant strains affect KdmB-mediated SMGC repression. For this, we analyzed emodin/monodictyphenone and F9775A/B (yellow polyketides), which are basically not produced in AnWT under our standard laboratory growth conditions (Cluster et al., 2010; Nützmann et al., 2011; Kleijnstrup et al., 2012). Consistent with our previous findings, transcription of *orsA* (encoding a non-reducing polyketide synthase (NR-PKS) required for orsellinic acid and F9775A/B biosynthesis) and *mdpG* (encoding a NR-PKS required for emodin biosynthesis) were strongly de-repressed in the cclAΔ strain resulting also in high levels of F9775A/B and emodin (Figure 1D). The metabolite levels in the sppAΔ strain, however, were almost unchanged; only a slight de-repression was evident at the transcriptional as well as at the metabolite level. Taken together, loss of CclA (but not SppA) largely phenocopied kdmBΔ with regard to secondary metabolism in *A. nidulans*.

Although we could not grow the setAΔ and H3K4R strains under same liquid AMM conditions, we were able to analyze their SM profiles from mycelia grown on solid ACM. We noticed that both strains were deeply pigmented when grown on this medium and subsequent mycelial extracts verified the accumulation of emodin and F9775A/B in both mutant strains, reminiscent of the high metabolite levels in cclAΔ and kdmBΔ grown in liquid AMM (Supplementary Figure S4). Overall, it was surprising to observe that both the lack of H3K4 methylation (setAΔ, H3K4R, cclAΔ) and the lack of KdmB – leading to a global increase of H3K4 methylation – result in similar phenotypes, i.e., reduced
penicillin G and sterigmatocystin and at the same time strongly increased emodin and F9775A/B production.

**Histone Demethylase-Independent Role of KdmB in Secondary Metabolite Gene Regulation in A. nidulans**

Contrary to its homolog Jhd2 in *S. cerevisiae* KdmB harbors, in addition to the JmjC and JmjN domain, a second PHD-type zinc finger as well as a C5HC2-type zinc finger and an ARID/BRIGHT DNA-binding domain (Figure 2A). A similar domain structure is present in Lid2 and Lid in *Schizosaccharomyces pombe* and *D. melanogaster*, respectively. Similar to Lid, one of the PHD domains may interact with histones, but the functions and targets of other domains remain opaque (Li et al., 2010).

We wanted to define whether the demethylase activity of KdmB is required for its function in SM gene activation (PEN and ST clusters) or repression (MDP and ORS clusters). For this, we have generated a demethylase-deficient mutant by introducing two point mutations in the JmjC domain of KdmB.
that were previously shown in *D. melanogaster* to abolish Fe\(^{2+}\) binding in the catalytic site and consequently abrogate Lid demethylase function (Li et al., 2010). These iron-coordinating amino acids are conserved in the KdmB JmjC domain, and hence, histidine 643 (H643) and glutamic acid 645 (E645) in *F. graminearum* (C) The *kdmB* transcript levels of the *A. nidulans* wild-type strain (AnWT), *kdmB*Δ, KdmB*, and *kdmB*Δ/*kdmB*Δ strains were measured as an additional control of generated strains. Prior to RNA extraction, indicated strains were grown in liquid AMM for 48 h. Experiments were performed in duplicates, and *kdmB* expression of AnWT was arbitrarily set as 1. Mean values and standard deviations are shown. RE, relative expression (D) For radial growth analysis, indicated strains were grown on solid ACM and AMM plates at 37°C under dark conditions. Pictures were taken after 5 days. Experiments were done in triplicates. Mean values and standard deviations are shown (E) For western blot analysis, indicated strains were grown in liquid AMM with 10 mM sodium nitrate for 48 h prior to protein extraction. Subsequently, whole protein extracts were isolated from lyophilized mycelia, and roughly, 10 μg of proteins were used for western analysis. H3 C-Term, H3K4me2, and me3 antibodies were used for detection. A densitometric analysis was performed and the protein levels of AnWT were arbitrarily set as 1. Experiments were done three times giving the same trend. Hence, only one experiment is depicted here.

Several independent mutants with identical growth phenotypes were obtained in each case (Supplementary Figure S6), and subsequently, one of each was used for further analysis. RT-qPCR of *kdmB* transcript levels in both the KdmB* and the *kdmB*/kdmB\(^{\text{Cm}}\) verified wild type-like *kdmB* levels (Figure 2C). No significant alterations in radial hyphal growth were visible when the generated strains were grown on solid ACM or AMM (Figure 2D). Noteworthy, biomass was slightly but significantly increased to about 130% in case of both the KdmB* and the *kdmB*Δ strain when cultivated under SM-inducing conditions (liquid AMM with 10 mM sodium nitrate).
The complemented kdmBΔ/kdmBΔ* strain showed a wild-type-like growth phenotype (Supplementary Figure S7). Western blot analysis of kdmBΔ, KdmB*, and kdmBΔ/kdmBΔ* strains using anti-H3K4 methylation-specific antibodies (H3K4me2 and H3K4me3) revealed elevated H3K4me3 but reduced H3K4me2 levels in both kdmBΔ and KdmB* compared to AnWT and the complemented strain kdmBΔ/kdmBΔ*. This phenotype is in accordance with our previous proteomic analyses (Gacek-Matthews et al., 2016) and likely attributable to the demethylase deficiency in both strains (Figure 2E).

We used these strains now to gain a deeper insight into the relevance of the KdmB histone demethylase function for SM gene regulation and cultivated them for 48 h under SM-inducing conditions (AMM with 10 mM sodium nitrate). For subsequent SM quantification, we focused on those clusters that have been identified as being KdmB- and H3K4 methylation-dependent in their activation (penicillin G and sterigmatocystin), or repression (emodin and F9775A/B). Surprisingly, both activated clusters remained silent not only in the kdmBΔ mutant but also in the KdmB* strain indicating that the full integrity of the JmjC demethylase domain with iron-coordinating H642 and E644 residues is crucial for the activating function of KdmB (Figure 3A). However, in respect to the repressing function of KdmB, we found that only the repression of F9775A/B biosynthesis was dependent on these conserved amino acids. Emodin, on the contrary, was not affected. The strain harboring the mutated KdmB (KdmB*) produced very low levels of this metabolite and showed only background expression of the biosynthetic gene (mdpG), similar to the A. nidulans wild-type control (Figure 3B). Taken together, KdmB functions in gene induction (penicillin G and sterigmatocystin) as well as repression (F9775A/B) in a strictly histone demethylase-dependent manner, while the repression of emodin biosynthesis appears to depend on KdmB but is independent of its function as a histone demethylase.

The Histone Demethylase-Independent Role in Secondary Metabolite Gene Regulation Is Conserved Also in the Phytopathogenic Fungus Fusarium graminearum

KDM5 proteins belong to the diverse JmjC domain-containing superfamily, which have been identified in all living organisms from bacteria to higher eukaryotes and are characterized by the highly conserved JmjC domain (Accari and Fisher, 2015). Thus, we next analyzed whether the function of the histone demethylase and the histone demethylase-independent function of KdmB* is conserved in other filamentous fungi. For this, we identified and deleted as well as mutated the KDM5 homolog designated FgKdm5, in the wheat pathogen F. graminearum PH-1 wild-type strain (FgWT). Three independent Δfgkdm5 mutant strains were produced, and full deletion was verified by diagnostic PCR and Southern blot (Supplementary Figures S8A,B,C). Next to Δfgkdm5, the JmjC domain of FgKdm5 was mutated at equivalent positions like the A. nidulans KdmB*, i.e., histidine 641 was mutated to glycine (H641G) and glutamic acid 643 to glutamine (E643Q). Out of this, three independent FgKdm5H641G,E643Q strains were obtained, hereafter referred to as FgKdm5*. Successful generation of both mutant strains was subsequently verified by diagnostic PCR (Supplementary Figures S8D,E). In addition, the native FgKDM5 was reintroduced into Δfgkdm5 in order to complement the loss of FgKDM5. Notably, in situ integration was unsuccessful so far. Therefore, FgKDM5 was introduced heterologously at a random location in the genome (ex situ), but expression was still driven by its native promoter and terminator sequences. Obtained mutants designated Δfgkdm5/FgKDM5* (complementation ex situ) were verified by diagnostic PCR (Supplementary Figures S8F,G) and transcription of FgKDM5 tested by RT-qPCR. This analysis verified functional integration of the complementing gene as wild type-like FgKDM5 levels were detected in both FgKdm5* and Δfgkdm5/FgKDM5*, while the transcript was absent from the Δfgkdm5 strain (Supplementary Figure S9A). Western blot analysis using anti-H3K4 methylation-specific antibodies for mono-, di-, and trimethylation confirmed the non-functional demethylase domain of FgKdm5* as we observed increased H3K4 tri- and reduced dimethylation levels in the mutant protein extracts compared to FgWT (Supplementary Figure S9B). H3K4 di- and trimethylation levels were restored to FgWT levels in the complemented Δfgkdm5/FgKDM5* strain (Supplementary Figure S9B).

Similar to the observations in A. nidulans, hyphal growth of Δfgkdm5 and FgKdm5* was not significantly altered when grown on solid Fusarium minimal or complete medium, FMM and FCM, respectively (Figure 4A and Supplementary Figure S9C). We also tested the involvement of FgKdm5 and its demethylase domain in the production of SMs. It is well known that F. graminearum sports a great variety of these compounds. 67 putative gene clusters have been characterized to be involved in SM biosynthesis (Sieber et al., 2014). Next to aurofusarin (Kim et al., 2005; Malz et al., 2005), deoxynivalenol (Kogel et al., 2005; Life, 2006; Audenaert et al., 2013), zearalenone (Lysoe et al., 2006), and fusaricin C (Farber and Sanders, 1986), several novel products as fusarielins (Sørensen et al., 2014; Wollenberg et al., 2016), fusaristatins (Sørensen et al., 2014), and chrysogine (Wollenberg et al., 2017) have been characterized in F. graminearum. To analyze changes in secondary metabolism, generated strains were grown together with FgWT for 2 weeks on potato dextrose agar (PDA) as described by Giese et al. (2013). Similar to A. nidulans and F. fujikuroi (Gacek-Matthews et al., 2016; Janevská et al., 2018), deletion of FgKDM5 resulted in a significant decrease of several SMs (Figure 4B). More specifically, production of the mycotoxins deoxynivalenol and fusarin C, the mycoestrogens zearalenone and fusarielin H as well as production of chrysogine was significantly decreased in strains lacking FgKdm5, while aurofusarin and fusaristatin A levels remained unaffected (Figure 4C and Supplementary Figure S10). To our surprise, most of the SMs affected by Δfgkdm5 appeared to be regulated by FgKdm5 independently of the integrity of the JmjC domain and its putative histone demethylase function as deoxynivalenol, zearalenone, fusarin C, and fusarielin H production levels are reaching FgWT levels in the FgKdm5* strain (Figure 4C). The only SM analyzed here which biosynthesis required the Fe³⁺-coordinating residues
FIGURE 3 | KdmB has a histone demethylase-independent role in regulation of the emodin gene cluster. (A,B) For SM analysis, the A. nidulans wild-type strain (AnWT), kdmBΔ, KdmB*, and kdmBΔ/kdmBΔ strains were grown in liquid AMM with 10 mM sodium nitrate for 48 h. Supernatants were applied for HPLC-MS/MS measurements. To exclude falsifications in SM biosynthesis due to differences in hyphal growth, SM production was correlated to biomass formation. For a direct comparison, production of the wild type was arbitrarily set to 100%. For statistical analysis, a student’s t test was performed: *p < 0.05; **p < 0.01; n.d., not detected. For determination of transcript levels, RNA was extracted of 48 h old, lyophilized mycelia. The transcription levels of ipnA [encoding the non-ribosomal peptide synthetase (NRPS) required for penicillin (PEN) production], aflR [encoding the pathway-specific transcription factor for sterigmatocystin (ST) biosynthesis], orsA [encoding a non-reducing polyketide synthase (NR-PKS) required for orsellinic acid and F9775A/B biosynthesis], and mdpG (encoding a NR-PKS required for emodin biosynthesis) were determined via RT-qPCR. SM data represent metabolite accumulation over time, whereas gene transcription was measured at a definite time point (ipnA, aflR, mdpG: 48 h; orsA: 24 h). Expression level of AnWT was arbitrarily set as 1 for the individual SM genes. Experiments were done in triplicates. Mean values and standard deviations are shown. RE, relative expression.
was chrysogine. Chrysogine was decreased in both, ∆fgkdm5 and FgKdm5* strains (Figure 4C). As expected, production level of aurofusarin and fusaristatin A that were not affected in ∆fgkdm5 remained unchanged also in FgKdm5* (Supplementary Figure S10). Complementation of ∆fgkdm5 with the native FgKDM5, i.e., ∆fgkdm5/FgKDM5Ces restored productions levels to FgWT (Figure 4C). Taken together, several SMs in F. graminearum depend on FgKdm5 but unlike KdmB in A. nidulans, the JmjC domain of FgKdm5 is not crucial for this activation function. Only in case of chrysogine, the histone demethylase function of FgKdm5 is essential.

kdmB Cannot Rescue the Demethylase-Dependent Phenotype in ∆fgkdm5

Next, we analyzed if kdmB of A. nidulans mediates activation of SMs in F. graminearum and if the integrity of the KdmB
JmjC domain is necessary for this activation function. To test this, the native kdmB was amplified from A. nidulans genomic DNA and sequence-verified plasmids integrated into the native FgKDM5 locus in F. graminearum. This resulted in Δfgkdm5/kdmBCA (complementation in situ) in which kdmB is driven by the F. graminearum promoter (Supplementary Figures S11A,B). Subsequent RT-qPCR of the kdmB transcript in three independent Δfgkdm5/kdmBCA mutants verified expression of kdmB (Supplementary Figure S11C). As expected and similar to Δfgkdm5, the Δfgkdm5/kdmBCA strains did not show an altered growth phenotype compared to FgWT when grown on solid FCM or FMM (Supplementary Figures S12A,B). To test whether kdmB can complement for the loss of FgKDM5 with regard to secondary metabolism, FgWT, Δfgkdm5, and Δfgkdm5/kdmBCA strains were grown for up to 2 weeks on PDA. The Δfgkdm5/kdmBCA strain rescued SM levels in case of deoxynivalenol, zearalenone, fusaric acid, fusarilin H, and chrysogine like in the Δfgkdm5 strain (Figure 5A). It is noteworthy that chrysogine appeared to be the only SM analyzed here that strictly depended on a functional histone demethylase domain (Figure 5B), suggesting that kdmB cannot rescue the histone demethylase function in Δfgkdm5. Intrigued by these findings, we next performed western blot experiments using anti-H3K4me3-specific antibodies. Indeed, H3K4me3 levels are similarly increased in Δfgkdm5/kdmBCA as compared to Δfgkdm5, while the complementation with the native gene (Δfgkdm5/FgKDM5CA) resulted in H3K4me3 that were comparable to FgWT (Figure 5C). Thus, cross-complementation experiments where Δfgkdm5 is complemented with kdmB suggest that kdmB can complement the histone demethylase-independent functions of FgKdm5 in SM gene regulation but not JmjC-dependent functions.

DISCUSSION

H3K4me3 is a well-studied chromatin mark involved not only in transcriptional initiation and elongation but also in recombination site selection during meiosis (Acquaviva et al., 2013; Adam et al., 2018). H3K4me3 are found predominantly at transcriptional start sites of transcribed genes (Santos-Rosa et al., 2002; Soares et al., 2017). Consequently, proteins involved in writing and erasing this histone PTM are associated with transcriptional activation and repression, respectively. H3K4me3 is established and erased by members of COMPASS/Set1 and KDM5, respectively. The latter belong to the Jumonji domain proteins of the KDM5 family and are involved in recognition and removal of H3K4me3 marks and thus resulting typically in repression of the targeted loci (Klose et al., 2006, 2007; Seward et al., 2007; Thynes et al., 2014). However, these proteins consist of several domains that might be important for their diverse functions in chromatin and transcriptional regulation.

Previously we could show that KdmB is mediating not only transcriptional silencing but also transcriptional activation of SMGCs in A. nidulans (Gacek-Matthews et al., 2016). In detail, KdmB serves as a canonical, transcriptional repressor for the
SMGCs monodictyphenone/emodin, and F9775A/B in *A. nidulans*. Strikingly, our data show that only for mediating repression of the F9775A/B gene cluster the integrity of the JmjC domain is important. Contrary to this, emodin biosynthesis and *mdlpG* expression did not depend on a functional JmjC domain as KdmB* mostly phenocopied AnWT. Demethylase-independent functions of KDM5 proteins have been described, but mostly in context with gene activation. For example, studies in *D. melanogaster* showed that loss of the essential KDM5 homolog Lid (little *imaginal discs*) results in retarded larval development (Li et al., 2010), and this delay is independent of the histone demethylase activity encoded by the JmjC domain, suggesting that other domains of Lid are critical during early developmental processes (Drelon et al., 2018). Furthermore, functions in gene regulation that are independent of the catalytic activity are not restricted to KDM5 homologs or histone demethylases. Thus, demethylase-independent regulations have also been described for the Lysine Specific Demethylase 1 LSD1 in human cancer cells. LSD1 promotes the survival of prostate cancer cells independently of its demethylase function (Sehrawat et al., 2018). Also, the methyltransferase Set1 in *S. cerevisiae* exhibits chromatin regulatory functions independent of H3K4 methylation (Lee et al., 2018). Here, the enzymatic activity of Set1 might be directed to a non-H3K4 substrate, which in turn has chromatin-associated regulatory functions (Mikhayeva et al., 2014). Thus, it might be possible that in our case KdmB utilizes a different domain to repress the emodin gene cluster in *A. nidulans*. However, strains largely lacking H3K4 tri-, di-, and monomethylation (setΔA, H3K4R and cclΔA) as well as loss of KdmB show overlapping phenotypes with regard to emodin and F9775A/B biosynthesis. Nevertheless, the increase of both SMs was still significantly lower in *kdmBΔ* as compared to cclΔA, suggesting that KdmB is part of a repression complex that requires H3K4me3 or me2 and that these marks are crucial for mediating repression. Whether the repressive function of KdmB is mediated directly at these loci and via H3K4me-binding or happens through the recruitment of different trans-factors (e.g., deacetylases) still remains to be elucidated. So far, we could not find any repressing function of FgKdm5 in *F. graminearum* regarding the SMs analyzed in this study. This is in agreement with recently published data in *F. fujikuroi*, where all SMs analyzed were found to be decreased by deletion of *FFKDM5* (Janevska et al., 2018). However, a genome-wide transcriptional analysis in this fungus showed that FfKdm5 also functions in SM gene repression although to a lesser extent. Thus, it is likely that some SMGCs are repressed by FgKdm5 also in *F. graminearum* similar to *F. fujikuroi* and *A. nidulans*.

Activating functions of KdmB were found at the penicillin G and sterigmatocystin biosynthesis loci in *A. nidulans*. In these cases, production of both SMs strictly depends on a functional histone demethylase-domain of KdmB. Additionally, our data demonstrate that H3K4 methylation is needed at these gene loci for activation as cclΔA largely phenocopies the loss of KdmB. This SM-activating function is conserved also in *F. graminearum*. In fact, from the analyzed SMs that are affected by the loss of FgKdm5, all of them were found to be downregulated in Δfgkdm5. Similarly, lack of the KDM5 homolog KdmB in *Epichloë festucae* resulted in reduced expression of the SMGCs involved in lolitremer (*ltm*) as well as ergoalkaloïds and indole diterpenes (*eas*) biosynthesis (Lukito et al., 2019). Contrary to *A. nidulans* but in line with *F. fujikuroi* (Janevska et al., 2018), *Fusarium* strains deficient for H3K4 methylation did not phenocopy the Δfgkdm5 strain. More specifically, only production of the virulence factor deoxynivalenol was found to be regulated in a similar manner in Δfgkdm5 and Δfgccl1, while fusarin and zearalenone biosynthesis were regulated antagonistically in the two strains [this study; Studt et al., 2017]. It is noteworthy that the same goes true for *F. fujikuroi*. Here, Δffkdm5 and Δffccl1 both resulted in reduced levels of gibberellic acid (Studt et al., 2017; Janevska et al., 2018), the virulence factor that facilitates colonization of rice roots by this fungus (Wiemann et al., 2013). Aurofusarin biosynthesis remained unaffected in both Δfgkdm5 and Δfgccl1.

The obvious difference between cclΔA on the one hand and Δfgccl and Δffccl1 on the other hand is the remaining H3K4 methylation levels. While hardly any H3K4 mono-, di-, and trimethylation is detectable in cclΔA, only H3K4 trimethylation is affected in both Δfgccl and Δffccl1 (Studt et al., 2017). The latter being in agreement with strains deleted for cclA in *E. festucae* that showed reduced but not abolished H3K4me3 levels (Lukito et al., 2019). Here, the *LTM* and *EAS* cluster genes are silenced by CcIA. Notably, both SMGCs reside in subtelomeric regions. It is intriguing to speculate that the distinct H3K4 methylation levels account for the observed differences in the phenotypes with regard to *kdmBΔ* as well as Δfgkdm5 and Δffkdm5. Surprisingly and again contrary to our data in *A. nidulans*, FgKdm5 activates the majority of the analyzed SMGCs (deoxynivalenol, zearalenone, fusarin C, and fusarilin H) independently of the demethylase-domain. Only the SMGC involved in chrysogine biosynthesis seems to be dependent on the functional JmjC domain of FgKdm5. Thus, it is possible that the activating functions of KDM5-proteins are conserved between *A. nidulans* and *F. graminearum*, but different domains are utilized for this task. In respect to SM regulation, the activating H3K4me3 histone mark is largely absent from most of the SMGCs in *A. nidulans* (Gacek-Matthews et al., 2016) (Supplementary Figure S13). The same is true for *F. graminearum* (Connolly et al., 2013) and *F. fujikuroi* (Wiemann et al., 2013; Studt et al., 2017; Janevska et al., 2018). These data would be consistent with two interpretations of SMGC regulation by KDM5 family members: either KDM5 homologs function in regulating SMGCs indirectly via trans-acting factors and/or by utilizing different domains for activating or repressing targeted SMGCs. Next to JmjC, Kdm5 harbors five further putative domains: a JmjN (IPR003349), an AT-rich interaction domain (ARID, IPR001606), two plant homeodomain (PHD) – type zinc fingers (IPR001965) and a C5HC2-type zinc finger (IPR004198). Interestingly, similar KDM5-domain studies in *D. melanogaster* revealed that activation of essential mitochondrial genes happens independently of the JmjC demethylase activity but relies on the C-terminal PHD domain that binds to H3K4me2/3 (Liu and Secombe, 2015). Thus, it would be interesting to see whether one of the aforementioned domains is involved in direct targeting and/or binding.
Mutation or deletion of single domains will show if one of them is required for transcriptional regulation in the future.

Cross-complementation experiments revealed that KdmB could only partially complement the Δfgkdm5 phenotype. Global H3K4me3 levels were enriched in both Δfgkdm5/kdmBΔ and Δfgkdm5. In agreement with this, chrysogine levels – the only SM analyzed in F. graminearum that was shown to strictly depend on the FgKdm5 demethylase function – were similarly decreased in both Δfgkdm5/kdmBΔ and Δfgkdm5. By contrast, all SMs that are regulated independently of the demethylase function were fully restored to FgWT levels in a Δfgkdm5/kdmBΔ strain. Notably, although domain synteny is conserved, KdmB and FgKdm5 exhibited a rather low amino acid sequence identity of only 58%. Amino acid sequence analysis of the different domains revealed quite high identities for the JmjC (87%), ARID/Bright (86%), and the C-terminal PHD (82%) domain, whereas rather low identities were gained for the JmjN (69%), the N-terminal PHD (55%) domain, and the C5HC2 type zinc finger (41%). Thus, it is surprising that it is the catalytic function of the JmjC domain that is not rescued by KdmB in a Δfgkdm5 strain, while demethylase-unrelated phenotypes are restored in the cross-complementation approach with regard to SM gene regulation. Thus, it would be interesting to analyze whether KDM5 orthologs from other fungal species are able to functionally complement the histone demethylase domain in Δfgkdm5. In a similar study, several orthologs from different distantly related Ascomycetes, i.e., A. nidulans (Eurotiomycetes), Alternaria alternata (Dothideomycetes), F. fujikuroi (Sordariomycetes), and from the close relative Sclerotinia sclerotiorum (Leotiomycetes), failed to functionally replace the H3K36-specific demethylase (Δbcdkm1) in Botrytis cinerea (Schumacher et al., 2019). Similar to our study, the proteins exhibited overall low sequence conservation, e.g., the sequence identity between KdmA and BcKdm1 was only 31.1%. However, it remains to be seen whether this is due to a non-functioning histone demethylase as observed for KdmB expressed in the Δfgkdm5 deletion strain.

Overall, this study demonstrates the complex regulatory circuits of KDM5 homologs in the two Ascomycetes A. nidulans and F. graminearum. In both organisms, a functional homolog of KDM5, i.e., KdmB and FgKdm5, respectively, is required for wild type-like SM gene regulation. Interestingly, the integrity of the JmjC domain is not always essential for mediating repression or activation of specific SMGCs. At which loci, KdmB/FgKdm5 function mediated indirectly via targeting H3K4me3 mainly present outside SMGCs and/or directly by interacting with chromatin complexes present at the regulated SMGCs remains to be elucidated. Future ChIP-seq analyses using tagged KDM5 proteins will allow us to map their genomic positions under different metabolic conditions and thereby help us to understand their bipartite function in activation and repression of SMGCs mechanistically.

DATA AVAILABILITY

The datasets generated for this study are available on request to the corresponding author.

AUTHOR CONTRIBUTIONS

SB, LS, and JS contributed to the design of the work. SB, JLS, MS, and AG-M were involved in data acquisition. SB, LS, JS, MS, and AG-M were involved in data analysis. SB, LS, and JS wrote the manuscript. All authors revised and approved the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2019.01759/full#supplementary-material

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