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Published in: Fungal Genetics and Biology

DOI (link to publication from Publisher): 10.1016/j.fgb.2021.103602

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Publication date: 2021

Document Version Publisher's PDF, also known as Version of record

Link to publication from Aalborg University

Citation for published version (APA):

Bachleitner, S., Sulyok, M., Sørensen, J. L., Strauss, J., & Studt, L. (2021). The H4K20 methyltransferase Kmt5 is involved in secondary metabolism and stress response in phytopathogénic Fusarium species. Fungal Genetics and Biology, 155, Article 103602. https://doi.org/10.1016/j.fgb.2021.103602

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### Fungal Genetics and Biology

journal homepage: www.elsevier.com/locate/yfgbi





# The H4K20 methyltransferase Kmt5 is involved in secondary metabolism and stress response in phytopathogenic *Fusarium* species

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### ARTICLE INFO

# Keywords: Chromatin modification Histone methylation H4K20me3 Fusarium Stress response Secondary metabolism

#### ABSTRACT

Fusarium fujikuroi and Fusarium graminearum are agronomically important plant pathogens, both infecting important staple food plants and thus leading to huge economic losses worldwide. F. fujikuroi belongs to the Fusarium fujikuroi species complex (FFSC) and causes bakanae disease on rice, whereas F. graminearum, a member of the Fusarium graminearum species complex (FGSC), is the causal agent of Fusarium Head Blight (FHB) disease on wheat, barley and maize. In recent years, the importance of chromatin regulation became evident in the plant-pathogen interaction. Several processes, including posttranslational modifications of histones, have been described as regulators of virulence and the biosynthesis of secondary metabolites. In this study, we have functionally characterised methylation of lysine 20 histone 4 (H4K20me) in both Fusarium species. We identified the respective genes solely responsible for H4K20 mono-, di- and trimethylation in F. fujikuroi (FfKMT5) and F. graminearum (FgKMT5). We show that loss of Kmt5 affects colony growth in F. graminearum while this is not the case for F. fujikuroi. Similarly, FgKmt5 is required for full virulence in F. graminearum as  $\Delta fgkmt5$  is hypovirulent on wheat, whereas the F. fujikuroi  $\Delta ffkmt5$  strain did not deviate from the wild type during rice infection. Lack of Kmt5 had distinct effects on the secondary metabolism in both plant pathogens with the most pronounced effects on fusarin biosynthesis in F. fujikuroi and zearalenone biosynthesis in F. graminearum. Next to this, loss of Kmt5 resulted in an increased tolerance towards oxidative and osmotic stress in both species.

### 1. Introduction

Plant pathogenic fungi are the cause of severe diseases on agricultural crops and thus, lead to huge economic losses each year (Fausto et al., 2019). Apart from significant reduction in yield, the potential contamination of food and feed by potent mycotoxins accumulating during infection represents a severe problem (Fisher et al., 2012; Zain, 2011). The two Fusarium species Fusarium fujikuroi and Fusarium graminearum are the causal agents of bakanae disease of rice (Wiemann et al., 2013) and Fusarium Head Blight (FHB) disease of wheat and barley (Kazan et al., 2012), respectively, and they are occurring worldwide. Both fungi have developed highly efficient infection strategies coupled with the production of secondary metabolites (SMs), including potent mycotoxins to ensure a successful colonisation of the host (Brown et al., 2010; Goswami and Kistler, 2004; Munkvold, 2017;

Niehaus et al., 2017; Wiemann et al., 2013). As SM biosynthesis is energy-consuming, SM gene expression is tightly regulated and only initiated under certain conditions. Generally, genes involved in SM biosynthesis are physically linked and organised in clusters (Keller and Hohn, 1997), which facilitates co-regulation, for example by chromatin-based mechanisms (Collemare and Seidl, 2019; Pfannenstiel and Keller, 2019). Here, posttranslational modifications (PTMs) of histone proteins are well-known for controlling gene expression in response to environmental or developmental signals as they have the capacity to define the degree of compaction from an 'open' transcriptionally active euchromatin to 'silent' transcriptionally inactive heterochromatin (Bannister and Kouzarides, 2011). Thus, a certain combination of histone PTMs can strongly influence the accessibility of chromatin and further provide a recognition platform for other chromatin-modifying enzymes, remodelling complexes and transcription factors, which consequently regulate

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transcription of underlying genes (Jenuwein, 2002; Kouzarides, 2007; Wood et al., 2005). While many different histone PTMs are known to date, only a few have been characterised in detail thus far. Histone acetylation and methylation were the first PTMs to be discovered (Allfrey et al., 1964) and since then continue to be the most well characterised histone PTMs. Acetylation is associated with transcriptional activation, whereas methylation has a bivalent role in transcriptional activation and repression, depending on the modified histone residue as well as the degree of methylation. For example, methylation of histone 3 lysine 4 (H3K4) is a hallmark for transcriptionally active euchromatin (Li et al., 2008; Pekowska et al., 2011; Pinskaya and Morillon, 2009), while H3K27me3 and H3K9me3 are hallmarks of facultative and constitutive heterochromatin, respectively (Freitag, 2017; Ridenour et al., 2020; Selker, 2017). A less characterised histone PTM in fungi is H4K20 trimethylation that is considered a silencing mark in higher eukaryotes (Kourmouli et al., 2004; Schotta et al., 2004). Methylation of H4K20 is established by members of the KMT5 family, containing the catalytically conserved SET (Su(var)3-9, Enhancer-of-zeste and Trithorax) domain (Jones and Gelbart, 1993). In vertebrates KMT5 A (PR-Set7/Set8) establishes the mono-methylation, whereas the two enzymes KMT5 B and KMT5 C (Suv4-20 h1/2) catalyse di- and trimethylation of H4K20 (Fang et al., 2002; Nishioka et al., 2002; Schotta et al., 2004). In contrast, the fission yeast Schizosacharomyces pombe possesses only one KMT5 orthologue i.e., Set9, which is solely responsible for all three types of H4K20 methylation (Sanders et al., 2004). While H4K20 methylation has been associated with many biological functions in vertebrates (ensuring genome integrity (Beck et al., 2012; Li et al., 2016; Oda et al., 2009), DNA damage repair (H4K20me2) (Chitale and Richly, 2018; Paquin and Howlett, 2018) and chromatin compaction (H4K20me3) (Schotta et al., 2004; Sims et al., 2006; Wang and Jia, 2009)), studies in S. pombe demonstrated that H4K20 methylation does not affect gene expression or heterochromatin function (Sanders et al., 2004). Rather, it seems H4K20me is limited to a function in DNA damage response (Sanders et al., 2004; Wang et al., 2009). However, almost no knowledge exists about H4K20 methylation in filamentous fungi, except for Magnaporthe oryzae, where MoKmt5 was shown to be the sole histone methyltransferase involved in H4K20me3 (Pham et al., 2015).

In this study, we set out to get a better understanding on the role KMT5 homologues play in the two plant pathogens *F. fujikuroi* and *F. graminearum*. For this, the KMT5 homologues were identified and characterised with regard to growth, asexual development, stress response, secondary metabolism and virulence. The KMT5 homologues are solely responsible for all methylation degrees of H4K20me in both *Fusarium* spp. Loss of Kmt5 results in increased stress tolerance on media supplemented with oxidative and osmotic agents in both species, while overexpression of *KMT5* largely reversed this phenotype. Next to this, H4K20me is crucial for wild type-like secondary metabolism in both fungi. Notably, some observed phenotypes such as reduced hyphal growth and attenuated virulence were only attributed to one of the fusaria, suggesting additional species-specific functions of Kmt5.

### 2. Materials and methods

### 2.1. Fungal strains, media and growth conditions

The wild-type strains of *F. fujikuroi* (FfWT) IMI58289 (Commonwealth Mycological Institute, Kew, UK) and *F. graminearum* (FgWT) PH-1 (FGSC 9075, NRRL 31084) were used as parental strains for deletion as well as overexpression experiments. For protoplasting, DNA isolation and western blot analyses, all strains were grown in darkness for three days on solid complete media (CM) (Pontecorvo et al., 1953) covered with cellophane sheets (Folia Bringmann) at 30 °C and 20 °C in the case of *F. fujikuroi* and *F. graminearum*, respectively. Fungal growth tests were performed on complete (CM, potato dextrose agar PDA) and minimal (synthetic ICI) media (Geissman et al., 1966) for 5 days under dark

conditions at 30 °C and 20 °C for F. fujikuroi and F. graminearum, respectively. Additionally, fungal growth was monitored with a plate reader assay as described by Cánovas et al. (2017). Briefly, a 96-well plate containing 100 µL of desired solid medium (CM or ICI) was inoculated with 1,000 and 10,000 spores and incubated at 28  $^{\circ}\text{C}$  and 24  $^{\circ}\text{C}$  in the case of F. fujikuroi and F. graminearum, respectively. Absorbance was quantified at 595 nm every 60 min for 72 hrs in a synergy HT micro plate reader (Biotek). Medium inoculated with water only served as negative control, and was used to cancel background noises. The experiments were performed twice with technical triplicates for each strain and Fusarium spp. Fungal abiotic stress tests were performed on complete media supplemented with 1 M sorbitol and 1 M NaCl for osmotic stress; 5, 10 and 25 mM  $H_2O_2$  and 0.25 and 0.5 mM menadione-sodium bisulfite (MSB) for oxidative stress (Ram and Klis, 2006). Both growth and stress plate assays were inoculated either with agar plugs (5 mm in diameter) or with 1,000 spores. To account for stress-independent growth deficiencies, growth on supplemented media was related to the growth on the non-supplemented media in the respective strains.

For F. fujikuroi SM analysis, the respective strains were pre-grown for 72 h in 300 mL Erlenmeyer flasks with 100 mL Darken medium (DVK) (Darken et al., 1959) on a rotary shaker under dark conditions at 180 rpm and 30 °C. A 500 µL aliquot of this culture was used for inoculation of synthetic ICI (Imperial Chemical Industries, Ltd, UK) medium (Geissman et al., 1966) with 6 mM or 60 mM glutamine as sole nitrogen source. Incubation was carried out for additional 7 days prior to chemical analyses. For F. graminearum SM analysis, the respective strains were grown on PDA plates for 14 days in darkness at 20 °C for two weeks as described by Giese et al. (2013). For conidia production, F. fujikuroi strains were grown for 7 days on solid V8 medium (20% v/v vegetable juice, Campbell Food, Puurs, Belgium) at 20  $^{\circ}$ C and 12 h/12 h light-dark cycles. In the case of F. graminearum, the strains were grown in 50 mL mung bean broth in 250 mL baffled flasks at 20  $^{\circ}\text{C}$  and 140 rpm for three days under dark conditions to induce conidiation (Bai and Shaner, 1996).

### 2.2. Plasmid construction

Plasmids for F. fujikuroi and F. graminearum deletion, complementation and overexpression strains were generated using yeast recombinational cloning (Colot et al., 2006; 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 are displayed in Figure S2 A and Table S1. For deletion constructs, the upstream (5') and downstream (3') sequences of FfKMT5 and FgKMT5 were amplified from FfWT and FgWT genomic DNA, respectively. For this the following primer pairs were used: FfKmt5 5F and FfKmt5\_5R for upstream, FfKmt5\_3F and FfKmt5\_3R for downstream regions in the case of F. fujikuroi and FgKmt5 5F and FgKmt5 5R for upstream and FgKmt5 3F and Fg Kmt5 3R for downstream regions, respectively, in the case of *F. graminearum*. For both deletion constructs, hygromycin B was used for selection of positive transformants. The hph resistance cassette was amplified from pCSN44 (Staben et al., 1989) with the primer pair Hph\_F and Hph\_R. Complementation constructs were generated by amplifying the wild-type gene FfKMT5/FgKMT5 from parental genomic DNA. For both fusaria, FfKMT5/FgKMT5 are driven by the native promotor and amplified with the primer pairs FfKmt5\_5F and FfKmt5\_Cil\_1R in the case of F. fujikuroi and FgKmt5\_5F and FgKmt5\_ORF\_Tgluc\_R in the case of F. graminearum. Both genes, FfKMT5/FgKMT5, were fused to the glucanase terminator of Botrytis cinerea (BcTgluc) followed by the geneticin (G418) resistance cassette, genR, amplified from p\(Delta f g k d m 5 / F g K D M 5 \) (Bachleitner et al., 2019) using the primer pair Geni-F and Geni-Tgluc-R. Downstream sequences of *FfKMT5/FgKMT5* were amplified using the primer pairs FfKmt5\_3F// FfKmt5\_3R and FgKmt5\_3F//FgKmt5\_3R in the case of *F. fujikuroi* and *F.* gramineraum, respectively. Overexpression constructs were generated by amplifying the upstream (5') sequence of FfKMT5 and FgKMT5 from

FfWT and FgWT genomic DNA, respectively, with the following primer pairs: FfKmt5\_5F and YRC\_OE\_FfKmt5\_hph\_R in the case of F. fujikuroi and FgKmt5\_5F and YRC\_OE\_FgKmt5\_hph\_R in the case of F. graminearum. The upstream sequence was followed by the hph resistance cassette and the Aspergillus nidulans oliC promotor, which was amplified from pOE::KDM5 (Janevska et al., 2018b), with the primers TtrpC-hphR and pOliCR1 for both F. fujikuroi and F. graminearum. The constitutive oliC promoter was followed by the native KMT5 gene and downstream (3') sequence, amplified with the primers YRC\_OE\_pOliC\_FfKmt5\_F and FfKmt5\_3R from FfWT gDNA in the case of F. fujikuroi and YRC\_OE\_pOliC\_FgKmt5\_F and FgKmt5\_3R from FgWT gDNA in the case of F. graminearum, respectively. For yeast recombinational cloning, the Saccharomyces cerevisiae FY834 was transformed with the obtained fragments yielding pΔffkmt5, pΔfgkmt5, pFfKMT5<sup>Cis</sup>, pFgKMT5<sup>Cis</sup> (complementation in situ), pOE::FfKMT5 and pOE::FgKMT5 (overexpression). Correct assembly of the gained plasmids was verified by restriction digest and/or sequencing.

### 2.3. Fungal transformation

Transformation of *F. fujikuroi* IMI58289 was performed as described by Wiemann et al. (2013). For this, the deletion fragment was amplified from p $\Delta ffkmt5$  with the primer pair FfKmt5\_5F and Ff\_Kmt5\_3R using a proof-reading polymerase (Q5-polymerase, New England Biolabs). For complementation, roughly 10 µg of p*FfkMT5*<sup>Cis</sup> was digested with *BgI*I prior to transformation. In the case of *F. graminearum*, the split marker approach was used (Goswami, 2012). For this, the deletion fragment was amplified with primer pairs FgKmt5\_5F//Hph\_split\_F and FgKmt5\_3R//Hph\_split\_R from p $\Delta fgkmt5$ , whereas the complementation fragment was amplified with the primer pairs FgKmt5\_5F//Geni\_splitF and FgKmt5\_3R//Geni\_splitR from p $\Delta fgkmt5$ . Transformed protoplasts were regenerated as described by Studt et al. (2017). Positive transformants were selected on regeneration media containing 100 ppm (parts per million) of either hygromycin or geneticin.

In the case of Δffkmt5 and Δfgkmt5 mutants, homologous recombination events were verified with the primer pair dia\_FfKMT5\_F/ dia\_FgKMT5\_F and pCSN44\_trpC\_T for the upstream part and dia\_FfKMT5\_R/dia\_FgKMT5\_R and pCSN44\_trpC\_P2 for the downstream part (Fig. S2 A/B). Absence of the native wild-type gene was verified the primers dia\_FfKMT5\_WT\_F//dia\_FfKMT5\_WT\_R dia\_FgKmt5\_WT\_F//dia\_FgKmt5\_WT\_R in the case of F. fujikuroi and F. graminearum, respectively (Fig. S2 B). Accordingly, homologous integration of the complementation constructs was verified using the primer pairs dia\_FfKmt5\_3'//pKS-Gengpd\_P and dia \_FgKmt5\_R//pKS-Gengpd\_P (downstream region) for F. fujikuroi and F. graminearum, respectively. Presence of the native wild-type gene i.e., FfKMT5/ FgKMT5, was verified using the primers dia\_FfKMT5\_WT\_F// dia FfKMT5 WT R (FfKMT5) and dia FgKmt5 WT F//dia FgKmt5 WT R (FgKMT5) (Fig. S3 A). Absence of hph was verified by the inability to grow on CM plates supplemented with hygromycin B (data not shown). In the case of OE::FfKMT5 and OE::FgKMT5, homologous recombination events were verified with the primer pair dia\_FfKMT5\_F/dia\_FgKMT5\_F and pCSN44\_trpC\_T for the upstream part and dia\_FfKMT5\_R/ dia\_FgKMT5\_R and pCSN44\_trpC\_P2 for the downstream part. Presence of the native wild-type gene, FfKMT5/FgKMT5, was verified using the primers FfKMT5\_WT\_F//FfKMT5\_WT\_R (FfKMT5) and FgKmt5\_WT\_F// FgKmt5\_WT\_R (FgKMT5) (Fig. S3 B). Altogether, Δffkmt5\_T3,  $\Delta ffkmt5_T4$  and  $\Delta ffkmt5_T7$  as well as  $\Delta fgkmt5_T11$  and  $\Delta fgkmt5_T14$ , were obtained for F. fujikuroi and F. graminearum, respectively. All mutants of F. fujikuroi and F. graminearum showed an identical phenotype. Hence, Δffkmt5\_T3 and Δfgkmt5\_T11 were arbitrarily chosen for complementation approaches. This resulted in at least two independent mutants each for F. fujikuroi (Δffkmt5/FfKMT5<sup>Cis</sup>\_T1, T6, T9 and T14) and F. graminearum (Δfgkmt5/FgKMT5<sup>Cis</sup>\_T9 and T12), that showed identical phenotypes and correct in situ integration of FfKMT5 and FgKMT5, respectively (Fig. S3A). Overexpression of FfKMT5 and

FgKMT5 was achieved by exchanging the native promoters with the constitutive strong *oliC* promoter from *A. nidulans*. At least three independent mutants overexpressing FfKMT5 and FgKMT5 were obtained for *F. fujikuroi* (OE::FfKMT5\_T2, T3 and T11) and *F. graminearum* (OE:: FgKMT5\_T2, T10 and T11), respectively (Fig. S3B).

### 2.4. Standard molecular techniques

For DNA isolation, lyophilised mycelium was ground to a fine powder, re-suspended in extraction buffer and isolated as previously described (Cenis, 1992). Isolated genomic DNA was used for PCR amplification and Southern blot analysis. Deletion and complementation fragments were amplified with the proof-reading Q5 DNA Polymerase (New Englang Biolabs) and PCR reactions were set up according to the manufacturers' protocol. For diagnostic PCR, the GoTaq® Green Master Mix (Promega) was used and the PCR reactions were set up according to the users' manual. For Southern blot analyses, genomic DNA of FfWT/FgWT and the respective mutant strains i.e., Δffkmt5/Δfgkmt5 were digested with the enzyme XhoI. The digested DNA was separated on a 1% (w/v) agarose gel and subsequently transferred onto positively charged nylon membranes (Roche Diagnostics GmbH, Germany) by downward blotting. Probes were labelled with DIG-11-dUTP using the DIG-High Prime DNA Labelling and Detection Starter Kit II from Roche. In the case of F. fukikuroi, the downstream region, amplified with the primers FfKmt5 3F//FfKmt5 3R, was used for probing. In the case of *F*. graminearum, hph was used for probing and amplified with the primer pair Hph F//Hph R.

For expression analyses, RNA was extracted from lyophilised 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). 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 the KMT5 expression, the primers RT-qPCR\_FfKmt5\_F//R were used in the case of F. fujikuroi and RT-qPCR\_FgKmt5\_F//R in the case of F. graminearum, respectively (Table S1). The cDNA levels were related to the constitutively expressed reference genes actin (FFUJ\_02611, FGSG\_07335),  $\beta$ -tubulin (FFUJ\_07385, FGSG\_09530) and glyceraldehyde 3-phosphate dehydrogenase (FFUJ 13490, FGSG 06257). Used primers are listed in Table S1. Primer efficiencies in the RT-qPCR were kept between 90 and 110%. Relative expression levels were calculated using the  $\Delta\Delta$ Ct method (Pfaffl, 2001) and normalised to the respective wild type. Generated plasmids were extracted and purified from E. coli and S. cerevisiae with the GeneJET<sup>TM</sup> plasmid miniprep kit (Fermentas GmbH, St. Leon-Rot, Germany). For western blot analyses, mycelium from 3 days-old strains was ground to a fine powder and proteins were extracted as described (Studt et al., 2016). Depending on the antibody, 15-50 µg of proteins were used for SDS-Page and subsequent western blotting. The membrane was probed with 1:4,000 dilutions of anti-H3 C-Term (Active Motif, AM39451), anti-H4K20me3 (AM39671 and Abcam, ab9053), anti-H4K20me2 (AM35482) and anti-H4K20me1 (AM39728) primary antibodies and 1:10,000 dilutions of anti-mouse (Jackson, AB\_10015289) HRP conjugated secondary antibody. Chemiluminescence was detected with Clarity<sup>™</sup> ECL Western Substrate and ChemDoc™ XRS (Bio-Rad). Each western blot was performed at least two times for both fusaria.

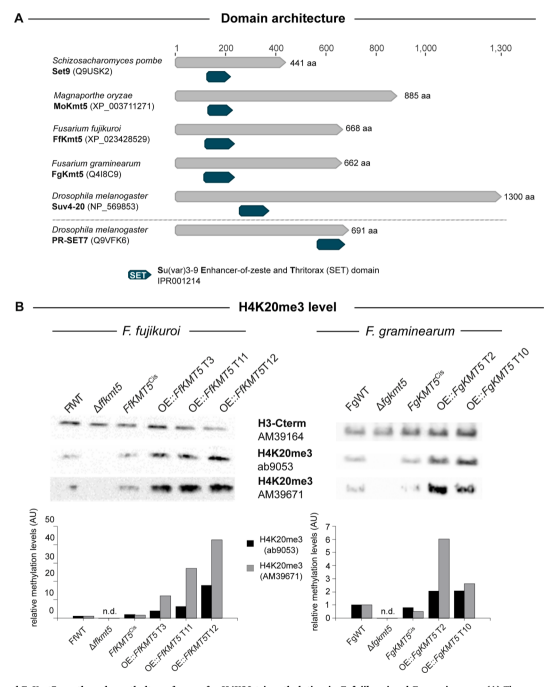
### 2.5. Pathogenicity assays

*F. fujikuroi* infection assays were performed according to Wiemann et al. (2013). In detail, five germinated rice seedlings (*Oryza sativa* sp. japonica cv. Nipponbare) were inoculated with each strain. In the case of  $\Delta ffkmt5$ ,  $\Delta ffkmt5/FfKMT5^{Cis}$  and OE::FfKMT5 at least two independent mutants were used. After 14 days at 28 °C, 80% humidity and a 12 h/12 h light–dark cycle, rice plants were screened for typical bakanae disease

symptoms and pictures were taken. Non-infected rice seedlings served as a negative control (MOCK). For *F. graminearum* infections the highly susceptible (FHB) USU-Apogee full-dwarf hard red spring wheat (*Triticum aestivum* cv. USU-Apogee; Reg.no CV-840, PI592742) was used. Infection experiments were carried out in biological triplicates and are described elsewhere (Studt et al., 2017). Briefly, for each biological replicate, two spikelets of five individual wheat plants were infected with 1000 spores each. As a control, ears were inoculated with sterile water only (MOCK). After each treatment, the ears were covered in

moistened plastic bags for the first 24 h to provide high humidity. Incubation conditions were set to 60% humidity, 20  $^{\circ}\text{C}$  for 16 h (day) and 18  $^{\circ}\text{C}$  for 8 h (night). The infection progress was monitored for up to 12 days.

In the case of *F. graminearum*, infection rate was determined as described elsewhere (Studt et al., 2017). Briefly, genomic DNA of infected plant material was extracted using the DNeasy® Plant Mini Kit (Qiagen). The infection rate was determined by qPCR quantification of the fungal DNA within the fungus/plant gDNA mixture. qPCR analysis



**Fig. 1. FfKmt5** and **FgKmt5** are the sole methyltransferases for H4K20 tri-methylation in *F. fujikuroi* and *F. graminearum*. (A) The conserved Kmt5 domain structure is shown for *Schizosaccharomyces pombe, Magnaporthe oryzae, F. fujikuroi* and *F. graminearum* as well as for the higher eukaryote *D. melanogaster*. In contrast to filamentous fungi and yeast, *D. melanogaster* harbours additionally to the H4K20 mono-methyltransferase PR-SET7, another enzyme i.e., Suv4-20 that is responsible for di- and tri-methylation of H4K20. Domain analysis was performed with protein sequences obtained from NCBI (accession number for each protein is shown in figure) in Geneious Prime 2020.2.4. (B) For western blot analysis, indicated strains were grown on complete media (CM) plates. Whole protein extracts were isolated from lyophilized mycelia and roughly 15 μg of proteins were used for western analysis. H3 C-Term and H4K20me3 antibodies were used for detection. Experiments were done in triplicates showing the same result. Hence, only one experiment is depicted here. For quantification a densitometric analysis was performed, and the respective wild-type strain was arbitrarily set to 1; AU, artificial units.

was performed using the iQTM SYBR Green Supermix (Bio-Rad) and primers Pks12\_ORF\_fwd//Pks12\_ORF\_rev for quantification of *F. graminearum* gDNA as well as primers ITS1P//ITS4 for quantification of plant gDNA.

### 2.6. Chemical analysis

Analysis of all here studied F. fujikuroi SMs was accomplished using the culture fluids of 7 days-old cultures that were directly used for analyses without further preparation. F. graminearum strains were grown for two weeks on potato dextrose agar (PDA) plates and subsequently extracted using MeOH/CH2Cl2/EtOAc (1/2/3, v/v), evaporated and resuspended in methanol/H<sub>2</sub>O (1/1, v/v) as previously described (Sondergaard et al., 2016). For quantification of GA<sub>3</sub> and DON levels of infected rice and wheat plants, freeze-dried and lyophilised plant material was pooled prior extraction (Studt et al., 2017). The samples were run on a QTrap 5500 LC-MS/MS System (Applied Biosystems, Foster City, CA, USA) equipped with a TurboIonSpray electrospray ionisation (ESI) source and a 1290 Series HPLC System (Agilent, Waldbronn, Germany). Chromatographic separation was done at 25 °C using a Gemini C18 150  $\times$  4.6 mm i.d., 5  $\mu$ m 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 (Sulyok et al., 2020).

### 3. Results

# 3.1. The histone methyltransferases FfKmt5 and FgKmt5 are solely responsible for mono-, di- and trimethylation of H4K20 in F. fujikuroi and F. graminearum, respectively

The KMT5 homologues in *F. fujikuroi* and *F. graminearum*, FfKmt5 and FgKmt5, respectively, were identified by determining the orthologues using QuartetS (Yu et al., 2011). The predicted proteins, FFUJ\_03480 (FfKmt5) and FGSG\_06529 (FgKmt5) in the case of *F. fujikuroi* and *F. graminearum*, respectively, are the orthologues of the H4K20-specific methyltransferase MoKmt5 (MGG\_07393) in *Magnaporthe oryzae* (Pham et al., 2015). Pairwise sequence alignment using LALIGN (Huang and Miller, 1991) showed 82.1% sequence identity (E-value 6.9e-172) of Kmt5 on protein level between the two fusaria, but only 48.5% (*F. fujikuroi*) and 48.2% (*F. graminearum*) sequence identity to *M. oryzae*. For illustration, all three sequences were aligned by T-coffee (Notredame et al., 2000) and formatted with BOXSHADE program (Fig. S1).

Similar to the homologue Set9 in *Schizosaccharomyces pombe* and MoKmt5 in *M. oryzae*, both FfKmt5 and FgKmt5 contain a catalytically active SET domain (Fig. 1 A). Contrary to vertebrates, fungi harbour only one KMT5 homologue, probably responsible for all three methylation degrees (Sanders et al., 2004) while higher eukaryotes harbour a H4K20 mono-methyltransferase (PR-Set7/Set8) (Fang et al., 2002; Nishioka et al., 2002) as well as one (*Drosophila melanogaster*) or more (vertebrates) di-and tri-methyltransferases (Suv4-20 enzymes) (Sakaguchi et al., 2008; Schotta et al., 2004). Notably, sequence analysis revealed that Set9 is more similar to Suv4-20 enzymes than to PR-SET7 (Greeson et al., 2008) (Fig. 1 A).

To study the functions of FfKmt5 and FgKmt5, the respective KMT5 open reading frames were deleted in the wild-type strains i.e., IMI58289 (F. fujikuroi) and PH-1 (F. graminearum), hereafter referred to as FfWT and FgWT, respectively (Fig. S2). Correct integration of the resistance cassettes was subsequently verified by diagnostic PCR and Southern blot analysis (Fig. S2 B/C). To test whether FfKmt5 and FgKmt5 are involved in writing H4K20 methylation in F. fujikuroi and F. graminearum, respectively, wild-type and the respective deletion strains were applied for western blotting using specific antibodies. All three methylation degrees of H4K20 were completely abolished in both  $\Delta ffkmt5$  and  $\Delta fgkmt5$  mutants (Fig. S5). To verify the obtained results, we next

approached complementation of Δffkmt5 and Δfgkmt5 as well as over-expression of KMT5 in both fusaria (Fig. S3). Notably, KMT5 transcript levels were restored to wild-type levels in the complemented strains, Δffkmt5/FfkMT5<sup>Cis</sup> and Δfgkmt5/FgkMT5<sup>Cis</sup> (hereafter referred to as FfkMT5<sup>Cis</sup> and FgkMT5<sup>Cis</sup>), and increased to about 10-fold in the over-expression strains, OE::FfkMT5 and OE::FgkMT5 when compared to the respective wild types (Fig. S4). Consistent with the expression data, H4K20me3 signals were wild type-like in the complemented strains and significantly increased in the OE::FfkMT5 and OE::FgkMT5 strains (Fig. 1B). Thus, Ffkmt5 and Fgkmt5 are solely responsible for writing mono-, di- and trimethylation of H4K20 in F. fujikuroi and F. graminearum, respectively.

### 3.2. FgKmt5 but not FfKmt5 significantly affects fungal colony growth

To gain insights into the importance of H4K20me for fungal growth, we monitored radial hyphal growth by performing plate assays using complete (CM and potato dextrose agar, PDA) as well as minimal (synthetic ICI medium supplemented with 6 mM glutamine as sole nitrogen source) media. To cover not only differences in fungal growth but also possible defects in germination, plates were inoculated with the relevant strains using agar plugs as well as conidia. No or only minor differences were detectable between \( \Delta ffkmt5 \) and \( \OE::FfKMT5 \) compared to FfWT, respectively, on both complete and minimal medium (Fig. 2A/B). In contrast to this, lack of FgKmt5 in F. graminearum resulted in significantly reduced radial growth on ICI and CM but not on PDA (Fig. 2A/B). Notably, the growth defect of  $\Delta fgkmt5$  was more obvious when inoculated with agar plugs, while conidia-inoculated plates converged to a wild type-like phenotype. Radial hyphal growth was reduced to 64% and 58% on ICI and CM, respectively, compared to FgWT when inoculated with agar plugs. However, when conidia were used as inoculum instead of agar plugs, the previously detected growth defect for \$\Delta fgkmt5\$ resembled a wild type-like phenotype. Here, only slight differences of growth were detected i.e., 75% and 90% on ICI and CM, respectively (Fig. 2A/B). As expected, complementation of  $\Delta fgkmt5$ i.e., FgKMT5<sup>Cis</sup>, restored the wild-type phenotype with regard to hyphal growth on all tested media. Notably, radial hyphal growth was also affected in OE::FgKMT5 strains although to a lesser extent. Here, radial hyphal growth was significantly reduced to 75% and 78% on ICI and PDA, respectively, compared to FgWT when agar plugs were used as inoculum. When conidia were used as inoculum instead of agar plugs, the previous detected growth defect for OE::FgKMT5 was still detectable on ICI (78% of FgWT) but not on PDA plates (Fig. 2A/B).

Radial hyphal extension (linear growth) does not necessarily reflect biomass accumulation. To evaluate whether biomass formation is indeed reduced in  $\Delta fgkmt5$  strains, we next performed plate reader assays with the *KMT5* deletion strains compared to both wild types. For this, absorbance (OD595 nm) was measured of fungal colonies growing on top of solid CM as well as on minimal medium (synthetic ICI with 6 mM glutamine as sole nitrogen source) as described (Cánovas et al., 2017). As expected, when grown on solid complete or minimal media, no differences in exponential growth were observed for  $\Delta ffkmt5$  compared to FfWT (Fig. S6). In agreement with the plate assay, biomass formation was reduced to 78% on minimal medium (synthetic ICI) in  $\Delta fgkmt5$  compared to FgWT, while no differences were observed when grown on solid CM (Fig. S6).

Next, we quantified spore formation in the two fusaria. For this, the respective strains were cultivated on solid V8 medium or in mung bean broth in the case of *F. fujikuroi* and *F. graminearum*, respectively. No significant differences in spore quantity were detectable for the *KMT5* deletion or overexpression strains when compared to the respective wild types in both fusaria (Fig. S7). This suggests that Kmt5 is not involved in asexual development in these fungi. Thus, deletion of *KMT5* impacts hyphal growth in *F. graminearum* but not in *F. fujikuroi*, while asexual development remains unaltered.

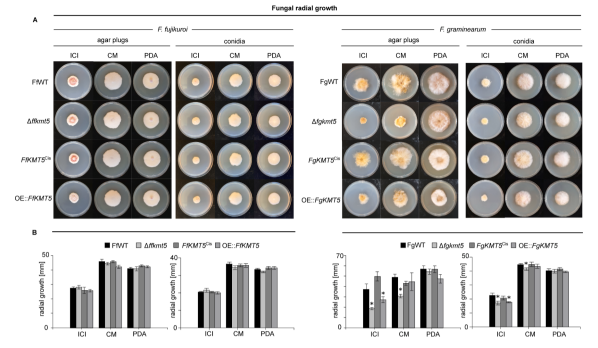


Fig. 2. Loss of Kmt5 results in impaired radial hyphal growth in *F. graminearum* but not in *F. fujikuroi*. (A) For growth analysis indicated strains were inoculated either with agar plugs or conidia on minimal media (ICI supplemented with 6 mM glutamine) and complete media (CM, PDA) and incubated at 30 °C and 20 °C for *F. fujikuroi* and *F. graminearum*, respectively. After five days post inoculation pictures were taken. (B) Experiments were performed in technical and biological triplicates. All experiments showed the same result, thus only one is depicted here. Mean values and standard deviations are shown, \*p < 0.05.

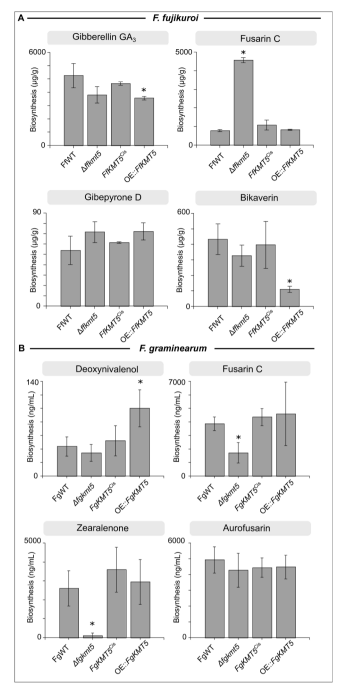
# 3.3. Both FfKmt5 and FgKmt5 are crucial for wild type-like secondary metabolism

Fusaria are known for their ability to produce a broad spectrum of SMs. Thus, we were interested to see if loss or overexpression of KMT5 affects SM biosynthesis in F. fujikuroi and F. graminearum. Both species produce sets of common but also species-specific SMs (Sieber et al., 2014; Wiemann et al., 2013). To analyse changes in secondary metabolism, F. fujikuroi strains were cultivated for seven days in synthetic ICI under nitrogen limited (6 mM glutamine) or nitrogen surplus (60 mM glutamine) conditions (Studt et al., 2017). F. graminearum strains were grown for two weeks on PDA plates prior to SM extraction as described by Giese et al. (2013). Overall, loss but also overexpression of KMT5 had distinct effects on some of the analysed SMs in both species. In detail, we observed small deviations in gibberellin (GA3) biosynthesis in F. fujikuroi. Here, overexpression but not deletion of FfKMT5 reduced GA3 levels to 67% of FfWT. It is noteworthy, that although  $\Delta$ ffkmt5 showed lower GA<sub>3</sub> levels, this trend was not significant. Similarly, gibepyrone D levels remained unaltered by loss or overexpression of FfKMT5. In contrast to GA3 and gibeyrone D levels, fusarin levels were strongly affected upon loss of FfKmt5 and increased to 585% when compared to FfWT. However, fusarin levels remained unaffected in OE::FfKMT5 strains. Further, bikaverin biosynthesis was significantly reduced to about 25% of FfWT in OE::FfKMT5 only (Fig. 3A). In F. graminearum, OE::FgKMT5 significantly increased DON levels to 229% of FgWT level, while DON biosynthesis remained wild type-like in  $\Delta fgkmt5$ . Contrary to F. fujikuroi, fusarin but also zearalenone levels were decreased to 43% and 4% compared to FgWT in  $\Delta fgkmt5$ , respectively. Biosynthesis of both compounds was not deviating from the wild type in OE::FgKMT5 strains. Aurofusarin biosynthesis remained unaffected by both,  $\Delta fgkmt5$ and OE::FgKMT5 (Fig. 3B). As expected, the complementation strains i. e., FfKMT5<sup>Cis</sup> and FgKMT5<sup>Cis</sup>, phenocopied the respective wild-type strains with regards to SM levels in both fusaria. Thus, altered KMT5 expression levels affected some SMs in both fusaria, with the most drastic phenotypes in fusarin biosynthesis in Δffkmt5 in F. fujikuroi and zearalenone biosynthesis in  $\Delta fgkmt5$  in F. graminearum. Notably,

deletion and respective overexpression strains did not show opposing phenotypes in any of the analysed *Fusarium* spp.

# 3.4. FgKmt5 but not FfKmt5 is required for wild type-like symptom development during pathogenic interaction

As overexpression of KMT5 affected biosynthesis of the virulence factors GA<sub>3</sub> and DON in axenic cultures in both fusaria, the role of Kmt5 in virulence and pathogenic development was investigated on rice (F. fujikuroi) and wheat heads (F. graminearum). In the case of F. fujikuroi, Δffkmt5, FfKMT5<sup>Cis</sup> and OE::FfKMT5 caused typical bakanae symptoms (shoot elongation and chlorosis) on rice seedlings, and internode elongation did not deviate from FfWT-infected rice seedlings (Fig. 4 A/B). The biosynthesis of GA3 is directly associated with the vellowish chlorotic leaves and hyper-elongated internodes of infected rice seedlings (Bömke and Tudzynski, 2009). Consistent with the occurrence of bakanae symptoms, GA3 levels quantified in planta were not affected in Δffkmt5 or OE::FfKMT5 when compared to FfWT (Fig. 4 C). The same is true for F. graminearum:  $\Delta fgkmt5$  or OE::FgKMT5 DON levels quantified in planta showed no deviations from both the FgWT and FgKMT5<sup>Cis</sup> (Fig. 4C), which contradicts axenic cultivations in the case of OE:: FgKMT5 (Fig. 3B). Yet, strains lacking FgKmt5 were hypovirulent on wheat (Fig. 4 A/B). While FgWT, FgKMT5<sup>Cis</sup> and OE::FgKMT5 were able to infect about 8 spikelets over a time period of 10 days, Δfgkmt5 was not able to infect more than 6 spikelets. Since experiments using biological systems may cause wide standard deviations, the experiment was carried out in three biological replicates. In all three experiments, the same trend was observed. In line with the hypovirulence, lower infection rates were determined for Δfgkmt5 i.e., 84% of FgWT, whereas complemented and overexpressed strains reached 90.5% and 95% (Fig. S8). Thus,  $\Delta fgkmt5$  is hypovirulent when compared to FgWT and FgKMT5<sup>Cis</sup>. Whether this phenotype is truly virulence-related or can be attributed to an impaired growth of  $\Delta \textit{fgkmt5}$  strains remains elusive at this point. To sum up, Kmt5 is not required for wild type-like virulence in F. fujikuroi, which is in line with previous observations in M. oryzae (Pham et al., 2015), but impacts virulence in *F. graminearum*.



**Fig. 3.** Loss of Kmt5 affects secondary metabolism in both fusaria. (A) For secondary metabolite analysis in *F. fujikuroi*, strains were cultured in liquid ICI (supplemented with 6 mM glutamine for giberellin, bikaverin and gibepyron, and 60 mM glutamine for fusarin C biosynthesis) for seven days at 30  $^{\circ}$ C and 180 rpm. (B) *F. graminearum* strains were grown on PDA plates for two weeks and subsequently extracted as described in Materials and Methods. Experiments were done in triplicates. Mean values and standard deviations are shown; \*p < 0.05.

# 3.5. Lack of Kmt5 results in mutants that are more tolerant to menadione sodium bisulfite-induced stress

Oxidative stress response protects organisms from harmful effects of reactive oxygen species (ROS), which can damage DNA, cellular components and cause disturbance of the cellular homeostasis (Angelova et al., 2005; Morano et al., 2012). H4K20me2 has been shown to be an important mark for genomic maintenance and DNA damage response

(Paquin and Howlett, 2018). Thus, we next analysed whether FfKmt5 and/or FgKmt5 plays a role in the cellular redox balance during the adaptation to environmental changes. For this, radial hyphal growth was investigated in the presence of two widely used agents to study oxidant stress i.e., hydrogen peroxide (H2O2) and menadione sodium bisulfite (MSB - a water-soluble derivate of menadione/vitamin K3). MSB has the ability to easily cross membranes and catalyses reactions, which release superoxide, hydrogen peroxide and hydroxyl radicals (Lehmann et al., 2012; Rasheed et al., 2018). KMT5 deletion, complementation and overexpression strains of both fusaria were grown on solid CM supplemented with either H2O2 or MSB. As differences of previous detected growth defects of  $\Delta fgkmt5$  were linked to the type of inoculation, we inoculated the relevant strains either with agar plugs or conidia. Furthermore, growth on media supplemented with stressors was normalised with normal growth on CM plates of the respective strain. After five days of fungal growth no deviations were detected for all strains on CM supplemented with H<sub>2</sub>O<sub>2</sub> when agar plugs were used (Fig. S9). In the case of conidia-inoculated plates, Δffkmt5 did not deviate from the wild type. However, FfKMT5<sup>Cis</sup> and OE::FfKMT5 showed retarded growth i.e., 80% and 73%, respectively, compared to FfWT on 5 mM H<sub>2</sub>O<sub>2</sub> plates. Similar to F. fujikuroi, no growth deviations were observed for \( \Delta fgkmt5 \) but slightly increased growth was detected for FgKMT5<sup>Cis</sup> and OE::FgKMT5 i.e., 116% and 113% when compared to FgWT on 0.5 mM H<sub>2</sub>O<sub>2</sub> and 115% and 107% on 1 mM H<sub>2</sub>O<sub>2</sub> plates, respectively (Fig. S9). As the complementation strains are not able to complement the wild-type phenotype in both fusaria, these data need to be evaluated carefully. It is noteworthy, that conidia of F. graminearum were not able to germinate on CM plates supplemented with 2.5 mM H<sub>2</sub>O<sub>2</sub> and higher. Thus, lower concentrations (0.5 and 1 mM H<sub>2</sub>O<sub>2</sub>) were chosen (Fig. S9).

While H<sub>2</sub>O<sub>2</sub> treatment did not have a compelling impact on the growth of the strains, a different scenario was shown when MSB was used as a stressor. Here, both  $\Delta ffkmt5$  and  $\Delta fgkmt5$  showed significantly increased growth in presence of MSB compared to the respective wildtype strains in the case of both inoculation methods (Fig. 5). In the case of F. fujikuroi, growth of \( \Delta ffkmt5 \) was increased to 205% of wildtype level on 0.5 mM MSB when inoculated with agar plugs, and 124% of wild-type level on 0.25 mM MSB plates when inoculated with conidia, respectively. Noteworthy, conidia were not able to germinate on 0.5 mM MSB plates. Thus, lower concentrations (0.1 and 0.25 mM MSB) were used. The complemented strain i.e., *FfKMT5*<sup>Cis</sup>, restored the wild type-growth phenotype in F. fujikuroi. Conversely, overexpression of FfKMT5 resulted in strains that are more susceptible towards MSB as radial growth rates were reduced to 60% on agar plug-inoculated MSB (0.5 mM) and 76% on conidia-inoculated MSB (0.25 mM) plates (Fig. 5A/B). In the case of F. graminearum, growth of  $\Delta fgkmt5$  was increased to 124% of wild-type level on agar plug-inoculated plates supplemented with 0.5 mM MSB, and to 164% of wild-type level on conidia-inoculated plates supplemented with 0.25 mM MSB. The  $\textit{FgKMT5}^{\text{Cis}}$  strain restored the wild-type phenotype. In contrast to F. fujikuroi, OE::FgKMT5 showed no consistent susceptibility towards MSB. Here, a reduced growth rate was only observed for conidia-inoculated plates supplemented with 0.1 mM MSB (Fig. 5A/B). To sum up, loss of Kmt5 resulted in elevated oxidative stress tolerance when treated with MSB in both fusaria. However, overexpression of KMT5 reversed the phenotype only in F. fujikuroi but not in F. graminearum.

### 3.6. Lack of Kmt5 results in mutants that are more tolerant to osmotic stress

Next, we analysed the adaptation ability to osmotic stress by growth on solid CM supplemented with high salt (NaCl) and sugar (sorbitol) concentrations. For this, the relevant strains were inoculated with either agar plugs or conidia together with the respective wild-type strains on solid CM supplemented with either 1 M NaCl or 1 M sorbitol. In both fusaria, lack of Kmt5 resulted in a significantly increased tolerance

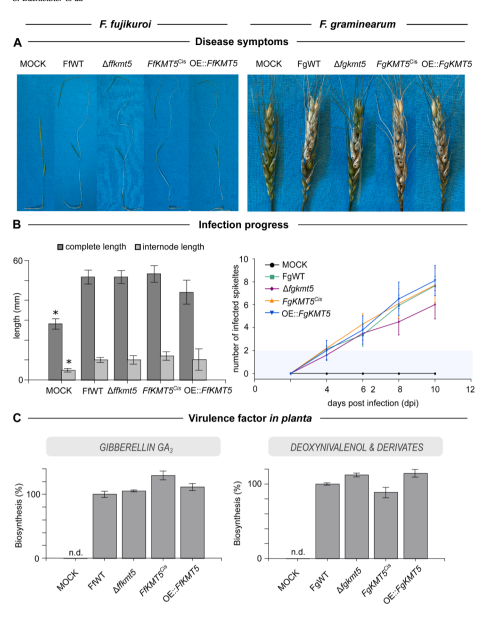


Fig. 4. Loss of FgKmt5 results in attenuated virulence on APOGEE wheat. (A) For pathogenicity analysis rice seedlings and wheat heads were infected with indicated F. fujikuroi and F. graminearum strains as described in Material and Methods. Pictures of rice plantlets and wheat heads were taken post-harvest. (B) Disease development was determined by measuring internode and complete length of rice plantlets in the case of F. fujikuroi infections. Wheat heads infected with F. graminearum strains were monitored over 10 days by measuring infected spikelets. Experiments were done in biological triplicates, resulting in the same trend, thus only one experiment is depicted here. (C) For in planta GA3 and DON quantification, five plants were infected with indicated F. fujikuroi and F. graminearum strains. Plants were pooled and the total amount of GA3 and DON was extracted and quantified by HPLC-MS/MS. Experiments were performed in biological triplicates; n.d., not detectable.

towards NaCl and sorbitol compared to the respective wild-type strains. In F. fujikuroi, radial growth of Δffkmt5 was increased to 118% in presence of 1 M NaCl and 112% in presence of 1 M sorbitol when agar plugs were used as inoculum. (Fig. 6). Similarly, increased osmotic stress tolerance was detected for conidia-inoculated plates. Here, radial growth of  $\Delta ffkmt5$  increased to 117% in presence of both, 1 M NaCl and 1 M sorbitol. The *FfKMT5*<sup>Cis</sup> strain restored the wild-type phenotypes for both inoculation methods. Conversely to \( \Delta ffkmt5, \) growth of OE:: FfKMT5 was slightly reduced on CM supplemented with 1 M sorbitol to 86% compared to FfWT, while no significant deviation was detected on NaCl supplemented plates (Fig. 6 A/B). A similar scenario was observed for F. graminearum. Here, growth of Δfgkmt5 was increased to 200% in presence of 1 M NaCl and 110% in presence of 1 M sorbitol when agar plugs were used as inoculum. Similarly, conidia-inoculated plates showed 110% and 116% increased growth for \( \Delta fgkmt5 \) in presence of 1 M NaCl and sorbitol, respectively. The complemented FgKMT5<sup>Cis</sup> strain restored only partially the wild-type phenotypes, as FgKMT5<sup>Cis</sup> was deviating from the respective wild type on agar-plug inoculated 1 M NaCl plates. In most cases, OE::FgKmt5 was not deviating from FgWT, except on agar plug-inoculated 1 M NaCl plates. Here, growth was similarly to FgKmt5<sup>Cis</sup> increased to 145% (Fig. 6A/B). Thus, loss of Kmt5

resulted in an increased tolerance towards osmotic stress in both fusaria, while only overexpression of *FfKMT5* reversed the phenotype on sorbitol-supplemented plates.

### 4. Discussion

Methylation of H4K20 is a well-studied histone PTM in animal cells and the fission yeast *Schizosaccharomyces pombe*. However, very little knowledge exists about its function in filamentous ascomycetes. In this study, we have identified and characterised the H4K20-specific methyltransferase Kmt5 in the two plant pathogenic fungi, *F. fujikuroi* and *F. graminearum*, by a reverse genetics approach.

# 4.1. Imbalanced H4K20me levels affect radial growth in F. graminearum but not in F. fujikuroi

While lack of *FfKMT5* did not affect morphology and growth in *F. fujikuroi*, we observed retarded growth for both the *FgKMT5* deletion and overexpression mutants on complete medium (agar plugs) and minimal media (ICI; conidia and agar plugs). This suggests that wild-type H4K20me levels are crucial for normal growth in *F. graminearum*.

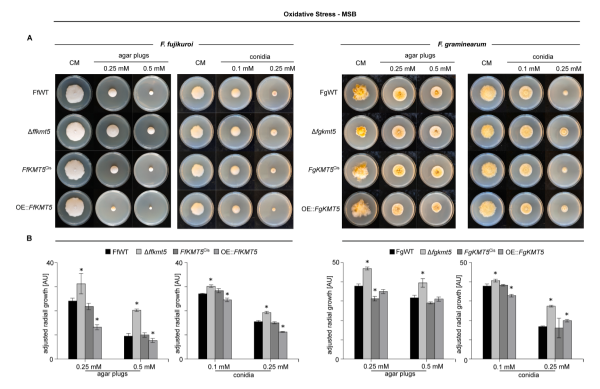


Fig. 5. Lack of Kmt5 results in strains more tolerant to menadione sodium bisulfite stress in F. fujikuroi and F. graminearum. (A) For oxidative stress analysis indicated strains were grown on complete media supplemented with 0.25 and 0.5 mM MSB when agar plugs were used as inoculum and 0.1 and 0.25 mM MSB when conidia were used for inoculation. Plates were cultivated at 30 °C and 20 °C in the case of F. fujikuroi and F. graminearum respectively. Pictures were taken after five days post inoculation. (B) Growth on media supplemented with stressors was adjusted to normal growth on CM plates of the respective strain. Experiments were performed in technical and biological triplicates. All experiments showed the same result, thus only one is depicted here. Mean values and standard deviations are shown; \*p < 0.05, AU; artificial units.

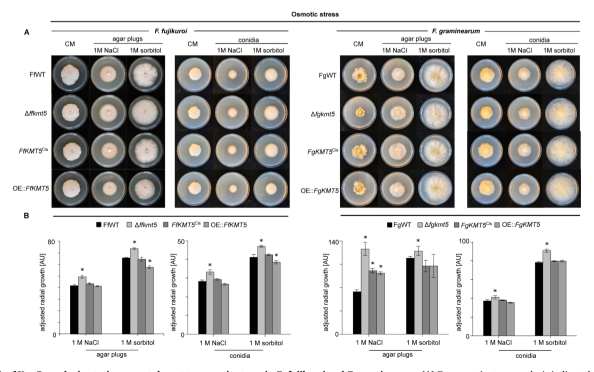


Fig. 6. Lack of Kmt5 results in strains more tolerant to osmotic stress in F. fujikuroi and F. graminearum. (A) For osmotic stress analysis indicated strains were grown on complete media supplemented with 1 M NaCl and 1 M sorbitol. Plates were inoculated either with agar plugs or conidia and incubated at 30 °C and 20 °C under dark conditions in the case of F. fujikuroi and F. graminearum respectively. Pictures were taken after five days post inoculation. (B) Growth on media supplemented with stressors was adjusted with normal growth on CM plates of the respective strain. Experiments were performed in technical and biological triplicates. All experiments showed the same result, thus only one is depicted here. Mean values and standard deviations are shown; \*p < 0.05., AU; artificial units.

Impaired growth was also reported for the rice blast fungus *M. oryzae*. Here, a lower growth rate was reported for Δ*mokmt5* compared to the wild-type strain on complete media (Pham et al., 2015). Loss of the Kmt5 orthologue Set9 was shown to be nonessential for cell viability in *S. pombe* but significantly affects DNA damage response (Sanders et al., 2004). In *D. melanogaster* but also in mammalian cells, loss of PR-Set7 disturbs the normal cell cycle progression and leads to accumulated DNA damage (Jørgensen et al., 2007; Li et al., 2016; Oda et al., 2009; Tardat et al., 2007). If imbalanced H4K20 methylation levels result in defective cell cycle progression and thus lead to growth impairment in *F. graminearum* remains unanswered at this moment. Similarly, why imbalanced H4K20 methylation levels only affect growth in *F. graminearum* but not in *F. fujikuroi* is elusive at this point. Characterisation of Kmt5 homologues in diverse fungal species will shed further light onto these species-specific differences.

### 4.2. Kmt5 affects SM biosynthesis in both fusaria

Deletion as well as overexpression of *KMT5* led to significant changes in SM biosynthesis in both Fusarium spp. In detail, GA<sub>3</sub> and bikaverin levels were significantly decreased by overexpression of FfKMT5 whereas fusarin levels were highly increased upon loss of FfKmt5 in F. fujikuroi. Thus, elevated H4K20me3 levels led presumably to a repression of these compounds and support the hypothesis of H4K20me3 as a silencing mark. However, a different scenario was shown for F. graminearum. Here, overexpression of FgKMT5 resulted in significantly increased DON levels, while loss of FgKmt5 significantly reduced and almost abolished biosynthesis of fusarins and zearalenone, respectively. This suggests activating functions of FgKmt5 for these SMs. As KMT5 deletion and overexpression strains did not show opposing phenotypes in any of the analysed Fusarium spp., we assume that alterations in SM biosynthesis are rather inflicted via secondary effects than by directly targeting SM biosynthetic genes. Notably, Xu and Kidder (2018) showed that H4K20me3 co-localises with H3K4me3 and H3K36me3 at transcriptionally dynamic regions in embryonic stem cells. Generally, H3K4me3 and H3K36me3 are considered hallmarks of active euchromatin (Black et al., 2012; Rivera et al., 2014), and many studies have highlighted their importance in SM regulation in filamentous fungi. For example, loss of Set1, the catalytic subunit of COMPASS, involved in H3K4 methylation reduced the expression of genes involved in DON and fumonisin biosynthesis in F. graminearum and F. verticilloides, respectively (Gu et al., 2017b; Liu et al., 2015). Similar results were obtained in Aspergillus flavus. Here, loss of Set1 led to abolished aflatoxin B1 biosynthesis (Liu et al., 2020). Notably, H3K4me3 was also shown to negatively impact SM biosynthesis. For example loss of another COM-PASS component (CclA) resulted in increased emodin and monodictyphenone levels in A. nidulans (Bachleitner et al., 2019; Bok et al., 2009). Similarly, in Aspergillus fumigatus and Aspergillus oryzae, loss of cclA increased the biosynthesis of gliotoxin and astellolides, respectively (Palmer et al., 2013; Shinohara et al., 2016). Furthermore, biosynthesis of five novel terpenoid compounds was induced in  $\Delta cclA$  mutants in Colletotrichum higginsianum (Dallery et al., 2019), and the subtelomeric gene clusters involved in ergotalkaloid (EAS) and lolitrem (LTM) biosynthesis were upregulated in Epichloë festucae (Lukito et al., 2019). In F. fujikuroi, loss of Set1 also resulted in increased production of bikaverin, fusarins and fusaric acid (Janevska et al., 2018b), thereby mirroring the  $\Delta ffkmt5$  phenotype. Similarly, deletion of CCL1 in F. fujikuroi and F. graminearum also resulted in an increased production of fusarins and bikaverin as well as zearalenone, respectively (Studt et al., 2017).

Less data is available for H3K36me3, though this histone PTM has been associated with SM biosynthesis in fusaria recently (Gu et al., 2017c; Janevska et al., 2018a). Notably, two proteins i.e., Set2 and Ash1, are associated with genome-wide H3K36me3 (Janevska et al., 2018a). While Set2 mediates H3K36 methylation in euchromatic regions, Ash1-mediated H3K36 methylation is predominately found in

subtelomeric regions in F. fujikuroi. Loss of either resulted in decreased  $GA_3$  but elevated fusarin and fusaric acid biosynthesis. Thus, strains deficient for H3K36me3 ( $\Delta set2$  and  $\Delta ash1$ ) also result in reduced  $GA_3$  levels but increased production of fusarins and fusaric acid in F. fujikuroi (Janevska et al., 2018a, 2018b; Studt et al., 2017), thereby again mirroring  $\Delta ffkmt5$  phenotype. Future research is required to elucidate the genome-wide positioning of H4K20me and to unravel its relationship with H3K4me3 and H3K36me3.

### 4.3. Lack of Kmt5 induces MSB- and osmotic stress tolerance in both fusaria

We found strains lacking Kmt5 more tolerant towards oxidative (MSB) stress in F. fujikuroi and F. graminearum. In the case of F. fujikuroi, overexpression of FfKmt5 reversed this phenotype on MSBsupplemented media resulting in a more sensitive phenotype. No growth changes were induced when H2O2 was used as an oxidative agent in both fusaria. Recently, Shao et al. (2019) showed that  $H_2O_2$  and MSB trigger different transcriptional pathways in A. oryzae. Generally, oxidative stress induces antioxidant enzymes including catalase (CAT), glutathione peroxidase (GPX) and superoxide dismutase (SOD) as well as the transcription factors Yap1 and Skn7 (Breitenbach et al., 2015; Lee et al., 1999; Morgan et al., 1997). In A. oryzae, different sets of CAT and SOD genes are activated by either H<sub>2</sub>O<sub>2</sub> or MSB (Shao et al., 2019). Further, MSB treatment increased GPX expression whereas H<sub>2</sub>O<sub>2</sub> rather triggered Yap1 transcription factors. Notably, different Skn7 transcription factors responded specifically either to H<sub>2</sub>O<sub>2</sub> or MSB induced stress (Shao et al., 2019). Thus, the observed diverging tolerance induced by H<sub>2</sub>O<sub>2</sub> and MSB stress of our KMT5 mutants might be owned from the different transcriptional response of oxidatively regulated genes. In addition to the increased MSB tolerance, lack of KMT5 resulted also in a higher tolerance to osmotic stress (NaCl and sorbitol) in both fusaria. In F. fujikuroi, overexpression of KMT5 reversed this phenotype resulting in strains that are more sensitive towards the osmotic stress (sorbitol). If the increased stress tolerance of the KMT5 deletion mutants results from alterations in DNA damage repair or through deregulation from stressassociated genes originally silenced by Kmt5 remains to be determined. The Hog1 pathway responds to osmotic stress but also contributes substantially to oxidative defence (de la Torre-Ruiz et al., 2015), and loss of the orthologue Hog1 i.e., FfSak1 and FgOS-2 resulted in mutants more sensitive to osmotic stress in F. fujikuroi and F. graminearum, respectively (Van Nguyen et al., 2012; Zheng et al., 2012). Thus, it is tempting to speculate that genes in this signal cascade are affected by Kmt5. A crosstalk between H4K20 and H3K9 methylation was shown to be not only fundamental for pericentric heterochromatin formation but also for DNA damage repair in higher eukaryotes (Kovaríková et al., 2018; Schotta et al., 2004). Schotta et al. (2004) demonstrated that pericentric H4K20me3 is severely reduced in Su(var)3-9 mutants. Further, studies in mammalian cells suggest a co-regulatory function between H3K9me3, H4K20me3 and 53BP1 (Kovaríková et al., 2018). It remains to be determined whether a similar crosstalk is happening also in Fusarium, but it is noteworthy that loss of the H3K9 methyltransferase Dim5 led to an increased tolerance towards osmotic stress in F. verticillioides, a phenotype which is attributed to a hyper-phosphorylation of Hog1 in  $\Delta fvdim5$  mutants (Gu et al., 2017a).

### 5. Conclusion

In this study, we have identified the histone methyltransferase Kmt5 that is solely responsible for mono-, di- and trimethylation of H4K20 in the two plant-pathogenic fungi *F. fujikuroi* and *F. graminearum*. Phenotypic characterisation of deletion as well as overexpression strains showed that Kmt5 is not essential for development and pathogenicity in both fusaria. While production of some SMs in *F. fujikuroi* and *F. graminearum* appear to rely on H4K20 methylation levels, others remain unaffected. Notably, observed findings are more likely to result from

secondary effects and not by Kmt5 directly targeting SM genes. Most strikingly, loss of Kmt5 resulted in increased tolerance to MSB and osmotic (NaCl and sorbitol) stress in both fusaria. Thus, Kmt5 is largely dispensable for basic metabolic and developmental processes in *F. fujikuroi* and *F. graminearum* but involved in MSB and osmotic stress response. Future studies on the genome-wide positioning of H4K20me3 will allow us to shed further light on the importance of this histone PTM in *Fusarium* spp.

### **Funding**

This work was funded by the FWF – Austrian Science Fund (Lise Meitner Grant M 2149-B22 to LS, Chrocosom P32790 to JS), and SB is the recipient of a PhD fellowship provided by NFB – NÖ Forschung und Bildung (SC16-026).

### CRediT authorship contribution statement

Simone Bachleitner: Methodology, Investigation, Validation, Formal analysis, Visualization, Writing - original draft. Michael Sulyok: Investigation, Resources, Writing - review & editing. Jens Laurids Sørensen: Investigation, Resources, Writing - review & editing. Joseph Strauss: Resources, Writing - review & editing, Supervision, Project administration, Funding acquisition. Lena Studt: Conceptualization, Methodology, Resources, Investigation, Supervision, Writing - original draft, Project administration, Funding acquisition.

### **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

### Appendix A. Supplementary material

Supplementary data to this article can be found online at https://doi.org/10.1016/j.fgb.2021.103602.

### References

- Angelova, M.B., Pashova, S.B., Spasova, B.K., Vassilev, S.V., Slokoska, L.S., 2005. Oxidative stress response of filamentous fungi induced by hydrogen peroxide and paraquat. Mycol. Res. 109, 150–158. https://doi.org/10.1017/ S0953756204001352.
- Allfrey, V.G., Faulkner, R., Mirsky, A.E., 1964. Acetylation and methylation of histones and their possible role in the regulation of RNA synthesis. Proc. Natl. Acad. Sci. USA 51, 786–794. https://doi.org/10.1073/pnas.51.5.786.
- Bachleitner, S., Sørensen, J.L., Gacek-Matthews, A., Sulyok, M., Studt, L., Strauss, J., 2019. Evidence of a demethylase-independent role for the H3K4-specific histone demethylases in Aspergillus nidulans and Fusarium graminearum secondary metabolism. Front. Microbiol. 10, 1759. https://doi.org/10.3389/ fmicb.2019.01759.
- Bai, G.H., Shaner, G., 1996. Variation in Fusarium graminearum and cultivar resistance to wheat scab. Plant Dis. 80, 975–979. https://doi.org/10.1094/pd-80-0975.
- Bannister, A.J., Kouzarides, T., 2011. Regulation of chromatin by histone modifications. Cell Res. 21, 381–395. https://doi.org/10.1038/cr.2011.22.
- Beck, D.B., Oda, H., Shen, S.S., Reinberg, D., 2012. PR-set7 and H4K20me1: at the crossroads of genome integrity, cell cycle, chromosome condensation, and transcription. Genes Dev. 26, 325–337. https://doi.org/10.1101/gad.177444.111.
- Black, J.C., Van Rechem, C., Whetstine, J.R., 2012. Histone lysine methylation dynamics: establishment, regulation, and biological impact. Mol. Cell 48, 491–507. https://doi. org/10.1016/j.molcel.2012.11.006.
- Bok, J.W., Chiang, Y., Szewczyk, E., Reyes-Domingez, Y., Davidson, A.D., Sanchez, J.F., Lo, H., Watanabe, K., Strauss, J., Oakley, B., Wang, C., Keller, N., et al., 2009. Chromatin-level regulation of biosynthetic gene clusters. Nat. Chem. Biol. 5, 462–464. https://doi.org/10.1038/nchembio.177.
- Bömke, C., Tudzynski, B., 2009. Diversity, regulation, and evolution of the gibberellin biosynthetic pathway in fungi compared to plants and bacteria. Phytochemistry 70, 1876–1893. https://doi.org/10.1016/j.phytochem.2009.05.020.
- Breitenbach, M., Weber, M., Rinnerthaler, M., Karl, T., Breitenbach-Koller, Lore, 2015. Oxidative stress in fungi: its function in signal transduction, Interaction Plant Hosts Lignocellulose Degradation. Biomolecules 5, 318–342. https://doi.org/10.3390/ biom5020318.

- Brown, N.A., Urban, M., van de Meene, A.M.L., Hammond-Kosack, K.E., 2010. The infection biology of Fusarium graminearum: Defining the pathways of spikelet to spikelet colonisation in wheat ears. Fungal Biol. 114, 555–571. https://doi.org/ 10.1016/j.funbio.2010.04.006.
- Cánovas, D., Studt, L., Marcos, A.T., Strauss, J., 2017. High-throughput format for the phenotyping of fungi on solid substrates. Sci. Rep. 7, 4289. https://doi.org/10.1038/ s41598-017-03598-9.
- Cenis, J.L., 1992. Rapid extraction of fungal DNA for PCR amplification. Nucleic Acids Res. 20, 2380. https://doi.org/10.1093/nar/20.9.2380.
- Chitale, S., Richly, H., 2018. H4k20me2: Orchestrating the recruitment of DNA repair factors in nucleotide excision repair. Nucleus 9, 212–215. https://doi.org/10.1080/ 19491034.2018.1444327.
- Collemare, J., Seidl, M.F., 2019. Chromatin-dependent regulation of secondary metabolite biosynthesis in fungi: Is the picture complete? FEMS Microbiology Reviews 43, 591–607. https://doi.org/10.1093/femsre/fuz018.
- Colot, H.V., Park, G., Turner, G.E., Ringelberg, C., Crew, C.M., Litvinkova, L., Weiss, R., Borkovich, K., Dunlap, J., et al., 2006. A high-throughput gene knockout procedure for *Neurospora* reveals functions for multiple transcription factors. Proc. Natl. Acad. Sci. 103, 10352–10357. https://doi.org/10.1073/pnas.0601456103.
- Dallery, J.F., Adelin, É., Le Goff, G., Pigné, S., Auger, A., Ouazzani, J., O'Connell, R., 2019. H3K4 trimethylation by CclA regulates pathogenicity and the production of three families of terpenoid secondary metabolites in *Colletotrichum higginsianum*. Mol. Plant Pathol. 20, 831–842. https://doi.org/10.1111/mpp.12795.
- Darken, M.A., Jensen, A.L., Shu, P., 1959. Production of gibberellic acid by fermentation. Appl. Microbiol. 7, 301–303. https://doi.org/10.1128/aem.7.5.301-303.1959.
- de la Torre-Ruiz, M., Pujol, N., Sundaran, V., 2015. Coping With Oxidative Stress. The Yeast Model. Curr. Drug Targets. 16, 2–12. https://doi.org/10.2174/ 1389450115666141020160105.
- Fang, J., Feng, Q., Ketel, C.S., Wang, H., Cao, R., Xia, L., Erdjument-Bromage, H., Tempst, P., Simon, J., Zhang, Y., 2002. Purification and functional characterization of SET8, a nucleosomal histone H4-lysine 20-specific methyltransferase. Curr. Biol. 12, 1086–1099. https://doi.org/10.1016/S0960-9822(02)00924-7.
- Fausto, A., Rodrigues, M.L., Coelho, C., 2019. The still underestimated problem of fungal diseases worldwide. Front. Microbiol. 10, 214. https://doi.org/10.3389/ fmicb.2019.00214.
- Fisher, M.C., Henk, D.A., Briggs, C.J., Brownstein, J.S., Madoff, L.C., McCraw, S.L., Gurr, S., 2012. Emerging fungal threats to animal, plant and ecosystem health. Nature 484, 186–194. https://doi.org/10.1038/nature10947.
- Freitag, M., 2017. Histone methylation by SET domain proteins in fungi. Annu. Rev. Microbiol. 71, 413–439. https://doi.org/10.1146/annurev-micro-102215-095757.
- Geissman, T.A., Verbiscar, A.J., Phinney, B.O., Cragg, G., 1966. Studies on the biosynthesis of gibberellins from (-)-kaurenoic acid in cultures of *Gibberella fujikuroi*. Phytochemistry 5, 933–947. https://doi.org/10.1016/S0031-9422(00)82790-9.
- Giese, H., Sondergaard, T.E., Sørensen, J.L., 2013. The AreA transcription factor in Fusarium graminearum regulates the use of some nonpreferred nitrogen sources and secondary metabolite production. Fungal Biol 117, 814–821. https://doi.org/10.1016/j.funbio.2013.10.006.
- Goswami, R.S., 2012. Targeted gene replacement in fungi using a split-marker approach. Methods Mol. Biol. 835, 255–269. https://doi.org/10.1007/978-1-61779-501-5\_16. Goswami, R.S., Kistler, H.C., 2004. Heading for disaster: Fusarium graminearum on cereal
- Goswami, R.S., Kistler, H.C., 2004. Heading for disaster: Fusarium graminearum on cerea crops. Mol. Plant Pathol. 5, 515–525. https://doi.org/10.1111/J.1364-3703.2004.00252.X.
- Greeson, N.T., Sengupta, R., Arida, A.R., Jenuwein, T., Sanders, S.L., 2008. Di-methyl H4 lysine 20 targets the checkpoint protein Crb2 to sites of DNA damage. J. Biol. Chem. 283, 33168–33174. https://doi.org/10.1074/jbc.M806857200.
- Gu, Q., Ji, T., Sun, X., Huang, H., Zhang, H., Lu, X., Wu, L., Huo, R., Wu, H., Gao, X., 2017a. Histone H3 lysine 9 methyltransferase FvDim5 regulates fungal development, pathogenicity and osmotic stress responses in *Fusarium verticillioides*. FEMS Microbiol Lett. 364, fxn184. https://doi.org/10.1093/femsle/fnx184.
- Gu, Q., Tahir, H.A.S., Zhang, H., Huang, H., Ji, T., Sun, X., Wu, L., Wu, H., Gao, X., 2017b. Involvement of FvSet1 in Fumonisin B1 Biosynthesis, Vegetative Growth, Fungal Virulence, and Environmental Stress Responses in Fusarium verticillioides. Toxins 9, 43. https://doi.org/10.3390/toxins9020043.
- Gu, Q., Wang, Z., Sun, X., Ji, T., Huang, H., Yang, Y., Zhang, H., Tahir, H.A.S., Wu, L., Wu, H., Gao, X., 2017c. FvSet2 regulates fungal growth, pathogenicity, and secondary metabolism in *Fusarium verticillioides*. Fungal Genet Biol. 107, 24–30. https://doi.org/10.1016/j.fgb.2017.07.007.
- Huang, X., Miller, W., 1991. A time-efficient, linear-space local similarity algorithm. Adv. Appl. Math. 12, 337–357. https://doi.org/10.1016/0196-8858(91)90017-D.
- Janevska, S., Baumann, L., Sieber, C.M.K., Münsterkötter, M., Ulrich, J., Kämper, J., Güldener, U., Tudzynski, B., 2018a. Elucidation of the Two H3K36me3 Histone Methyltransferases Set2 and Ash1 in Fusarium fujikuroi Unravels Their Different Chromosomal Targets and a Major Impact of Ash1 on Genome Stability. Genetics 208, 153–171. https://doi.org/10.1534/genetics.117.1119.
- Janevska, S., Güldener, U., Sulyok, M., Tudzynski, B., Studt, L., 2018b. Set1 and Kdm5 are antagonists for H3K4 methylation and regulators of the major conidiation-specific transcription factor gene ABA1 in Fusarium fujikuroi. Environ. Microbiol. 20, 3343–3362. https://doi.org/10.1111/1462-2920.14339.
- Jenuwein, T., 2002. Translating the Histone Code. Science 293, 1074–1080. https://doi. org/10.1126/science.1063127.
- Jones, R.S., Gelbart, W.M., 1993. The *Drosophila* Polycomb-group gene Enhancer of zeste contains a region with sequence similarity to trithorax. Mol. Cell. Biol. 13, 6357–6366. https://doi.org/10.1128/mcb.13.10.6357-6366.1993.
- Jørgensen, S., Elvers, I., Trelle, M.B., Menzel, T., Eskildsen, M., Jensen, O.N., Helleday, T., Helin, K., Sørensen, C.S., 2007. The histone methyltransferase SET8 is

- required for S-phase progression. J. Cell Biol. 179, 1337–1345. https://doi.org/
- Kazan, K., Gardiner, D.M., Manners, J.M., 2012. On the trail of a cereal killer: recent advances in *Fusarium graminearum* pathogenomics and host resistance. Mol. Plant Pathol. 13, 399–413. https://doi.org/10.1111/j.1364-3703.2011.00762.x.
- Keller, N.P., Hohn, T.M., 1997. Metabolic pathway gene clusters in filamentous fungi. Fungal Genet. Biol. 21, 17–29. https://doi.org/10.1006/fgbi.1997.0970.
- Kourmouli, N., Jeppesen, P., Mahadevhaiah, S., Burgoyne, P., Wu, R., Gilbert, D.M., Bongiorni, S., Prantera, G., Fanti, L., Pimpinelli, S., Shi, W., Fundele, R., Singh, P.B., 2004. Heterochromatin and tri-methylated lysine 20 of histone H4 in animals. J. Cell Sci. 117, 2491–2501. https://doi.org/10.1242/jcs.01238.
- Kouzarides, T., 2007. Chromatin modifications and their function. Cell 128, 693–705. https://doi.org/10.1016/j.cell.2007.02.005.
- Kovaríková, A.S., Legartová, S., Krejcí, J., Bártová, E., 2018. H3K9me3 and H4K20me3 represent the epigenetic landscape for 53BP1 binding to DNA lesions. Aging 10, 2585–2605. https://doi.org/10.18632/aging.101572.
- Lee, J., Godon, C., Lagniel, G., Spector, D., Garin, J., Labarre, J., Toledano, M.B., 1999.
  Yap1 and Skn7 control two specialized oxidative stress response regulons in yeast.
  J. Biol. Chem. 274, 16040–16046. https://doi.org/10.1074/jbc.274.23.16040.
- Lehmann, M., Laxa, M., Sweetlove, L.J., Fernie, A.R., Obata, T., 2012. Metabolic recovery of *Arabidopsis thaliana* roots following cessation of oxidative stress. Metabolomics 8, 143–153. https://doi.org/10.1007/s11306-011-0296-1.
- Li, F., Huarte, M., Zaratiegui, M., Vaughn, M.W., Shi, Y., Martienssen, R., Cande, W.Z., 2008. Lid2 is required for coordinating H3K4 and H3K9 methylation of heterochromatin and euchromatin. Cell 135, 272–283. https://doi.org/10.1016/j. cell.2008.08.036
- Li, Y., Armstrong, R.L., Duronio, R.J., Macalpine, D.M., 2016. Methylation of histone H4 lysine 20 by PR-Set7 ensures the integrity of late replicating sequence domains in *Drosophila*. Nucleic Acids Res 44, 7204–7218. https://doi.org/10.1093/nar/gkw333.
- Liu, Y., Liu, N., Yin, Y., Chen, Y., Jiang, J., Ma, Z., 2015. Histone H3K4 methylation regulates hyphal growth, secondary metabolism and multiple stress responses in Fusarium graminearum. Environ. Microbiol. 17, 4615–4630. https://doi.org/ 10.1111/1462-2920.12993.
- Liu, Y., Zhang, M., Xie, R., Zhang, F., Wang, S., Pan, X., et al., 2020. The methyltransferase AflSet1 is involved in fungal morphogenesis, AFB1 biosynthesis, and virulence of Aspergillus flavus. Front. Microbiol. 11, 234. https://doi.org/ 10.3389/fmicb.2020.00234.
- Lukito, Y., Chujo, T., Hale, T.K., Mace, W., Johnson, L.J., Scott, B., 2019. Regulation of subtelomeric fungal secondary metabolite genes by H3K4me3 regulators CclA and KdmB. Mol. Microbiol. 112, 837–853. https://doi.org/10.1111/mmi.14320.
- Morano, K.A., Grant, C.M., Moye-Rowley, W.S., 2012. The response to heat shock and oxidative stress in *Saccharomyces cerevisiae*. Genetics 190, 1157–1195. https://doi. org/10.1534/genetics.111.128033.
- Morgan, B.A., Banks, G.R., Mark Toone, W., Raitt, D., Kuge, S., Johnston, L.H., 1997. The Skn7 response regulator controls gene expression in the oxidative stress response of the budding yeast Saccharomyces cerevisiae. EMBO J 16, 1035–1044. https://doi.org/ 10.1093/emboi/16.5.1035.
- Munkvold, G.P., 2017. *Fusarium* species and their associated mycotoxins. Methods Mol. Biol. 1542, 51–106. https://doi.org/10.1007/978-1-4939-6707-0\_4.
- Niehaus, E.M., Kim, H.K., Münsterkötter, M., Janevska, S., Arndt, B., Kalinina, S.A., Houterman, P.M., Ahn, I.P., Alberti, I., Tonti, S., Kim, D., Sieber, C.M.K., Humpf, H., Yun, S., Güldener, U., Tudzynski, B., 2017. Comparative genomics of geographically distant Fusarium fujikuroi isolates revealed two distinct pathotypes correlating with secondary metabolite profiles. PLoS Pathog 13, e1006670. https://doi.org/10.1371/ journal.ppat.1006670.
- Nishioka, K., Rice, J.C., Sarma, K., Erdjument-Bromage, H., Werner, J., Wang, Y., Chuikov, S., Valenzuela, P., Tempst, P., Steward, R., Lis, J.T., Allis, C.D., Reinberg, D., 2002. PR-Set7 is a nucleosome-specific methyltransferase that modifies lysine 20 of histone H4 and is associated with silent Chromatin. Mol. Cell. 9, 1201–1213. https://doi.org/10.1016/S1097-2765(02)00548-8.
- Notredame, C., Higgins, D.G., Heringa, J., 2000. T-coffee: A novel method for fast and accurate multiple sequence alignment. J. Mol. Biol. 302, 205–217. https://doi.org/ 10.1006/jmbi.2000.4042
- Oda, H., Okamoto, I., Murphy, N., Chu, J., Price, S.M., Shen, M., Torres-Padilla, M., Heard, E., Reinberg, D., 2009. Monomethylation of histone H4-lysine 20 is involved in chromosome structure and stability and is essential for mouse development. Mol. Cell. Biol. 29, 2278–2295. https://doi.org/10.1128/mcb.01768-08.
- Palmer, J.M., Bok, J.W., Lee, S., Dagenais, T.R.T., Andes, D.R., Kontoyiannis, D.P., Keller, N., 2013. Loss of CclA, required for histone 3 lysine 4 methylation, decreases growth but increases secondary metabolite production in *Aspergillus fumigatus*. PeerJ 1, e4. https://doi.org/10.7717/peerj.4.
- Paquin, K.L., Howlett, N.G., 2018. Understanding the histone DNA repair code: H4K20me2 makes its mark. Mol. Cancer Res. 16, 1335-1345. https://doi.org/10.1158/1541-7786.MCR-17-0688.
- Pekowska, A., Benoukraf, T., Zacarias-Cabeza, J., Belhocine, M., Koch, F., Holota, H., Imbert, J., Andrau, J., Ferrier, P., Spicuglia, S., 2011. H3K4 tri-methylation provides an epigenetic signature of active enhancers. EMBO J 30, 4198–4210. https://doi. org/10.1038/emboj.2011.295
- Pfaffl, M.W., 2001. A new mathematical model for relative quantification in real-time RT-PCR. Nucleic Acid Res. 29 (9), e45. https://doi.org/10.1093/nar/29.9.e45.
- Pfannenstiel, B.T., Keller, N.P., 2019. On top of biosynthetic gene clusters: How epigenetic machinery influences secondary metabolism in fungi. Biotechnol. Adv. 37, 107345. https://doi.org/10.1016/j.biotechadv.2019.02.001.
- Pham, K.T.M., Inoue, Y., VuVan, B., Nguyen, H.H., Nakayashiki, T., Ikeda, K., Nakayashiki, H., 2015. MoSET1 (Histone H3K4 Methyltransferase in Magnaporthe

- Pinskaya, M., Morillon, A., 2009. Histone H3 lysine 4 di-methylation: A novel mark for transcriptional fidelity? Epigenetics 4, 302–306. https://doi.org/10.4161/
- Pontecorvo, G., Roper, J.A., Chemmons, L.M., Macdonald, K.D., Bufton, A.W.J., 1953. The genetics of *Aspergillus nidulans*. Adv. Genet. 5, 141–238. https://doi.org/10.1016/S0065-2660(08)60408-3.
- Ram, A.F.J., Klis, F.M., 2006. Identification of fungal cell wall mutants using susceptibility assays based on Calcofluor white and Congo red. Nat. Protoc. 1, 2253–2256. https://doi.org/10.1038/nprot.2006.397.
- Rasheed, R., Arslan Ashraf, M., Kamran, S., Iqbal, M., Hussain, I., 2018. Menadione sodium bisulphite mediated growth, secondary metabolism, nutrient uptake and oxidative defense in okra (Abelmoschus esculentus Moench) under cadmium stress. J. Hazard. Mater. 360, 604–614. https://doi.org/10.1016/j.jhazmat.2018.08.043.
- Ridenour, J.B., Möller, M., Freitag, M., 2020. Polycomb repression without bristles: facultative heterochromatin and genome stability in fungi. Genes (Basel) 11, 638. https://doi.org/10.3390/genes11060638.
- Rivera, C., Gurard-Levin, Z.A., Almouzni, G., Loyola, A., 2014. Histone lysine methylation and chromatin replication. Biochim. Biophys. Acta - Gene Regul. Mech. 1839, 1433–1439. https://doi.org/10.1016/j.bbagrm.2014.03.009.
- Sakaguchi, A., Karachentsev, D., Seth-Pasricha, M., Druzhinina, M., Steward, R., 2008. Functional characterization of the *Drosophila Hmt4-20/Suv4-20 histone* methyltransferase. Genetics 179, 317–322. https://doi.org/10.1534/ genetics.108.087650.
- Sanders, S.L., Portoso, M., Mata, J., Bähler, J., Allshire, R.C., Kouzarides, T., 2004. Methylation of histone H4 lysine 20 controls recruitment of Crb2 to sites of DNA damage. Cell 119, 603–614. https://doi.org/10.1016/j.cell.2004.11.009.
- Schotta, G., Lachner, M., Sarma, K., Ebert, A., Sengupta, R., Reuter, G., Reinberg, D., Jenuwein, T., 2004. A silencing pathway to induce H3–K9 and H4–K20 trimethylation at constitutive heterochromatin. Genes Dev 18, 1251–1262. https://doi.org/10.1101/gad.300704.
- Schumacher, J., 2012. Tools for *Botrytis cinerea*: New expression vectors make the gray mold fungus more accessible to cell biology approaches. Fungal Genet. Biol. 49, 483–497. https://doi.org/10.1016/j.fgb.2012.03.005.
- Selker, E.U., 2017. H3K27 methylation: a promiscuous repressive chromatin mark. Curr. Opin. Genet. Dev. 43, 31–37. https://doi.org/10.1016/j.gde.2016.11.001.
- Shao, H., Tu, Y., Wang, Y., Jiang, C., Ma, L., Hu, Z., Wang, J., Zeng, B., He, B., 2019. Oxidative stress response of Aspergillus oryzae induced by hydrogen peroxide and menadione sodium bisulfite. Microorganisms 7, 225. https://doi.org/10.3390/ microorganisms/080225
- Shinohara, Y., Kawatani, M., Futamura, Y., Osada, H., Koyama, Y., 2016. An overproduction of astellolides induced by genetic disruption of chromatinremodeling factors in *Aspergillus oryzae*. J. Antibiot. (Tokyo) 69, 4–8. https://doi. org/10.1038/ia.2015.73.
- Sieber, C.M.K., Lee, W., Wong, P., Mu, M., Mewes, H., Schmeitzl, C., varga, E., Berthiller, F., Adam, G., Güldener, U., et al., 2014. The Fusarium graminearum genome reveals more secondary metabolite gene clusters and hints of horizontal gene transfer. PLoS One 9, e110311. https://doi.org/10.1371/journal.
- Sims, J.K., Houston, S.I., Magazinnik, T., Rice, J.C., 2006. A trans-tail histone code defined by monomethylated H4 Lys-20 and H3 Lys-9 demarcates distinct regions of silent chromatin. J. Biol. Chem. 281, 12760–12766. https://doi.org/10.1074/jbc. M513462200.
- Sondergaard, T.E., Fredborg, M., Oppenhagen Christensen, A.M., Damsgaard, S.K., Kramer, N.F., Giese, H., Sörensen, J.L., 2016. Fast screening of antibacterial compounds from Fusaria. Toxins 8, 1–9. https://doi.org/10.3390/toxins8120355.
- Staben, C., Jensen, B., Singer, M., Pollock, J., Schechtman, M., Kinsey, J., Selker, E., 1989. Use of a bacterial hygromycin B resistance gene as a dominant selectable marker in *Neurospora crassa* transformation. Fungal Genet. Rep. 36, 22. https://doi. org/10.4148/1941-4765.1519.
- Studt, L., Janevska, S., Arndt, B., Boedi, S., Sulyok, M., Humpf, H.U., Tudzynski, B., Strauss, J., 2017. Lack of the COMPASS component Ccl1 reduces H3K4 trimethylation levels and affects transcription of secondary metabolite genes in two plant-pathogenic Fusarium species. Front. Microbiol. 7, 1–17. https://doi.org/10.3389/fmicb.2016.02144.
- Studt, L., Rösler, S.M., Burkhardt, I., Arndt, B., Freitag, M., Humpf, H.U., Dickschat, J.S., Tudzynski, B., 2016. Knock-down of the methyltransferase Kmt6 relieves H3K27me3 and results in induction of cryptic and otherwise silent secondary metabolite gene clusters in Fusarium fujikuroi. Environ. Microbiol. 18, 4037–4054. https://doi.org/ 10.1111/1462-2920.13427.
- Sulyok, M., Stadler, D., Steiner, D., Krska, R., 2020. Validation of an LC-MS/MS-based dilute-and-shoot approach for the quantification of > 500 mycotoxins and other secondary metabolites in food crops: challenges and solutions. Anal. Bioanal. Chem. 412, 2607–2620. https://doi.org/10.1007/s00216-020-02489-9.
- Tardat, M., Murr, R., Herceg, Z., Sardet, C., Julien, E., 2007. PR-Set7-dependent lysine methylation ensures genome replication and stability through S phase. J. Cell Biol. 179, 1413–1426. https://doi.org/10.1083/jcb.200706179.
- Van Nguyen, T., Schöfer, W., Bormann, J., 2012. The stress-activated protein kinase FgOS-2 is a key regulator in the life cycle of the cereal pathogen *Fusarium graminearum*. Mol. Plant-Microbe Interact. 25, 1142–1156. https://doi.org/10.1094/ MPMI-02-12-0047-R.
- Wang, Y., Jia, S., 2009. Degrees make all the difference: The multifunctionality of histone H4 lysine 20 methylation. Epigenetics 4, 273–276. https://doi.org/10.4161/ epi.4.5.9212.

- Wang, Y., Reddy, B., Thompson, J., Wang, H., Nomaichi, K., Yates, J.R., et al., 2009. Regulation of Set9-Mediated H4K20 Methylation by a PWWP Domain Protein. Mol. Cell. 33, 428–473. https://doi.org/10.1016/j.molcel.2009.02.002.
- Wiemann, P., Sieber, C.M.K., von Bargen, K.W., Studt, L., Niehaus, E.M., Espino, J.J., Huß, K., Michielse, K.B., Albermann, S., Wagner, D., Bergner, S.V., Conolly, L.R., Fischer, A., Reuter, G., Kleigrewe, K., Bald, T., Wingfield, B.D., Ophir, R., Freeman, S., Hippler, M., Smith, K.M., Brown, D.W., Proctor, R.H., Münsterkötter, M., Freitag, M., Humpf, H.U., Güldener, U., Tudzynski, B., 2013. Deciphering the cryptic genome: genome-wide analyses of the rice pathogen Fusarium fujikuroi reveal complex regulation of secondary metabolism and novel metabolites. PLoS Pathog. 9, e1003475. https://doi.org/10.1371/journal.ppat.1003475.
- Wood, A., Schneider, J., Shilatifard, A., 2005. Cross-talking histones: implications for the regulation of gene expression and DNA repair. Biochem. Cell Biol. 83, 460–467. https://doi.org/10.1139/o05-116.
- Xu, J., Kidder, B.L., 2018. H4K20me3 co-localizes with activating histone modifications at transcriptionally dynamic regions in embryonic stem cells. BMC Genomics 19, 514. https://doi.org/10.1186/s12864-018-4886-4.
- Yu, C., Zavaljevski, N., Desai, V., Reifman, J., 2011. QuartetS: A fast and accurate algorithm for large-scale orthology detection. Nucleic Acid Res. 39, 1–10. https:// doi.org/10.1093/nar/gkr308.
- Zain, M.E., 2011. Impact of mycotoxins on humans and animals. J. Saudi Chem. Soc. 15, 129-144. https://doi.org/10.1016/j.jscs.2010.06.006.
- Zheng, D., Zhang, S., Zhou, X., Wang, C., Xiang, P., Zheng, Q., Xu, J.R., 2012. The FgHOG1 pathway regulates hyphal growth, stress responses, and plant infection in Fusarium graminearum. PLoS ONE 7, e49495. https://doi.org/10.1371/journal. pone.0049495.