The SWC4 subunit of the SWR1 chromatin remodeling complex is involved in varying virulence of Metarhizium brunneum isolates offering role of epigenetic regulation of pathogenicity

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ABSTRACT

The host–pathogen interaction is a multifactorial process subject to a co-evolutionary arms race consisting of rapid changes in both host and pathogen, controlled at the genetic and epigenetic levels. Previously, we showed intra-species variation in disease progression and pathogenicity in aphids for Metarhizium brunneum isolates MBK and MB7. Herein, we compared genomic, epigenetic, and metabolomic variations between these isolates and their effects on pathogenicity. Genomic variation could not completely explain the observed differences between the isolates. However, differential N6-adenine methylation (6 mA) and its correlation to reduced expression of the essential SWC4 subunit of SWR1 chromatin-remodelling complex (SWR1-C) led us to hypothesize a role for swc4 in the varying pathogenicity. Mutagenesis of the essential swc4 gene in MBKisolate resulted in reduction of secondary-metabolite (SM) secretion and impaired virulence in Galleria mellonella. Our results suggest the role of SWC4 in the regulation of SMs and the role of both SWC4 and SWR1-C in virulence of M. brunneum isolates. A better understanding of epigenetic regulation of SM production and secretion in entomopathogenic fungi may enable manipulation for better biocontrol performance, and expand possibilities for environmentally friendly pest control.

Introduction

The host–pathogen interaction is a multifactorial process subject to a co-evolutionary arms race [1]. This arms race consists of rapid changes in not only the genomes, but also the behaviour of both the host and the pathogen, as each responds to their partner’s attacks [2]. Entomopathogenic fungi (EPF) are an environmentally friendly alternative to synthetic pesticides, and major players in integrated pest management towards reducing the use of chemical insecticides [3]. Metarhizium spp. (Hypocreales: Clavicipitaceae) are among the most prevalent and best-studied EPF. Moreover, Metarhizium spp. are a model system for host–pathogen interactions and a resource for biotechnological tools [4–6].

The life cycle of hypocrealean fungi, such as Metarhizium, can be divided into two major stages: the pathogenic lifestyle stage, involving pathogenic interactions of the asexual conidial with invertebrate hosts, and the saprophytic lifestyle stage, involving proliferation in the rhizosphere and on soil organic matter as a saprophyte. On susceptible hosts, the pathogenic lifestyle of hypocrealean EPF consists of primary infection on the host cuticle (adhesion, germination), followed by later stages of infection (penetration, colonization, conidiosgenesis) [7]. Infection progression requires the pathogen to be adapted to, and compatible with the host, and to possess the genes, proteins and secondary metabolites (SMs) needed for successful infection [8]. Species of Metarhizium secrete SMs which assist the fungi to evade and weaken insect’s immune system, such as destruxins [9].

Variation in host susceptibility can be attributed to both fungal and host characteristics [1,10,11]. In fungi, the penetration process has been extensively studied and shown to include the production of a versatile set of peptides [12–15]. These peptides contribute to fungal infection by...
of cuticle degrading enzymes [12]. In many cases, the host is unable to resist mycosis once penetration of the cuticle occurs [13–15].

Fungal infection of insect hosts progresses rapidly: studies have found that within 24 h of inoculation, there are changes in host gene expression in response to germination on the outer cuticle [16]. The regulation of pathogen attack and host counterattack is largely uncharacterized. Whereas spontaneous mutations in virulence and resistance genes are a key driver of this co-evolutionary arms race, epigenetic changes provide a mechanism for more rapid, stable shifts in gene expression, allowing for faster adaptation to evolutionary innovation in the partner.

Studies have shown the importance of epigenetic regulation, particularly in asexual organisms [17,18]. Epigenetic marks such as DNA methylation and histone acetylation can provide infected hosts with resistance to a fungal pathogen that can be transferred to subsequent generations [19]. Studies in recent years have demonstrated the importance of histone modifications in host–pathogen interactions [6,20]. Furthermore, the epigenetic regulators themselves, such as methyltransferases, can be strongly manipulated during infection and in turn, affect a cascade of induction and suppression of genes involved in virulence and pathogenicity [21].

The most common epigenetic DNA modification in eukaryotes is 5-methylcytosine (m5C), which has been studied in a variety of models and conditions. This modification is known to occur across the genomes of organisms from several kingdoms and control various processes [22]. N6-adenine methylation (6 mA), a well-known DNA modification in bacterial genomes, was recently identified at various levels in eukaryotic genomes [23,24], and shown to occur mainly in expressed genes [23]. In bacteria, 6 mA plays an essential role in virulence and protection from foreign DNA [25]. However, its role and mode of action in eukaryotes have yet to be determined.

Variations in EPF species’ virulence to different hosts can be attributed to the fungi’s “toolkit” and the level of compatibility with the host, and later, to their ability to evade the host’s immune system. For successful infection, fungi secrete SMs, which are organic compounds comprising antimicrobial and immunosuppressant compounds [26]. These can cause mortality even when used as purified extracts [27] and are one of the main factors controlling fungal compatibility with certain hosts [28,29]. Many SM gene clusters have been putatively found in EPF genomes, but their regulation mechanisms and final products are unknown [29–31]. The role of chromatin-modification and remodelling complexes in controlling SM production and fungal virulence has been elucidated [32–34]. One of the most interesting chromatin-remodelling complexes, which has yet to be demonstrated as a SM regulator in fungi, is the SWR1 complex (SWR1-C), which is involved in the deposition of histone variant H2A.Z into the nucleosome [32]. Its wide presence in eukaryotes, from humans (as SRCAP) to yeast, demonstrates its essential nature, as it plays critical roles in DNA repair and transcription regulation [32,35,36].

The proper selection of fungal strain is crucial for the effectiveness of integrated pest management against the desired insect pest [28,37]. Different EPF species have varying degrees of virulence depending on the host being challenged, especially when specialist strains are used. However, generalist fungal strains may also perform differently on different hosts [13,14,38]. Recently, we revealed that not only inter-species, but also intra-specie variations in M. brunneum affect virulence ability [7]. In the present study, we further explored the basis for better performance of one isolate over the other in the two M. brunneum isolates MbK and Mb7. Here, we focused on the role of epigenetic regulation and demonstrate how epigenetic regulators affect pathogenicity in M. brunneum.

Materials and methods

Fungal growth and maintenance

Two selected isolates of M. brunneum, Mb7 and MbK, were used in this study [7]. Fungi were cultured on SDA (Difco) plates at 28 ± 0.5°C in a four-day incubator. Sporulation occurred mainly during the first 14 d of incubation.

Genomic DNA extraction from fungal material

Conidia were collected from SDA plates using a sterile loop and placed in 2-mL screw-cap tubes (for the extraction of fungal gDNA from infected Myzus persicae aphids: 7 DPI, aphids [7] were collected in a 2-mL screw-cap tube and a similar extraction was carried out). Samples were ground with metal beads in the presence of lysis buffer (Lucigen, USA) using a Geno/Grinder (SPEX SamplePrep, USA) at 6500 oscillations/min for 2 min. Then, the samples were subjected to gDNA extraction using MasterPure Yeast DNA Extraction Kit (Lucigen) according to the manufacturer’s instructions. Samples were analysed by Nanodrop (Thermo Fisher Scientific, USA) and separated on a 1% agarose gel in the presence of ethidium bromide. Samples were maintained at −20°C until used.
Next-generation sequencing and bioinformatics analysis

Genomic DNA samples extracted from Mb7 and MbK fungal isolates were further purified using gDNA clean & concentrator Kit (Zymo, USA) and served for library construction and Illumina 150- paired end next-generation sequencing (Sequencing core, University of Illinois, USA). Raw paired-reads were subjected to a filtering and cleaning procedure. The FASTX Toolkit (http://hannonlab.cshl.edu/fastx_toolkit/index.html, version 0.0.13.2) was used to trim read-end nucleotides with quality scores <30, using the fastq_quality_trimmer, and to remove reads with less than 70% base pairs with a quality score ≤ 30 using the Fastq_quality_filter. Clean reads (20.5 and 14.9 paired-end million reads for Mb7 and MbK, respectively) were mapped to the *Metarhizium brunneum* (GCA_013426205.1_ASM1342620v1) genome using the Burrows–Wheeler Aligner (BWA) software 0.7.12-r1039 [39]. The resulting mapping files were processed using SAMtools/Picard tool [http://broadinstitute.github.io/picard/, version 1.78 [40]] for adding read group information, sorting, marking duplicates and indexing. Then, the local realignment process for locally realigning reads was performed so that the number of mismatching bases is minimized across all the reads using the RealignerTargetCreator and IndelRealigner of the Genome Analysis Toolkit version v4.1.9.0 [GATK; http://www.broadinstitute.org/gatk/ [41]]. Finally, the variant calling procedure was performed using HaplotypeCaller of the GATK toolkit (https://gatk.broadinstitute.org/hc/en-us) developed by Broad Institute of MIT and Harvard (Cambridge, MA, USA). SnpEff tool [version 5.0d [42]] was used for annotations and predictions the effects of genetic variants based on the annotation downloaded from the NCBI database (https://ftp.ncbi.nlm.nih.gov/genomes/all/GCA_013426205.1_ASM1342620v1/GCA_013426205.1_ASM1342620v1_genomic.gff.gz). Clean paired-end reads were assembled de novo using SPAdes toolkit [v3.14.1 [43]]. Scaffold above 2000 bp were analysed for gene prediction using OmicsBox (v2.0.36) based on AUGUSTUS software [44]. The predicted proteins were used as a query term for a search of the NCBI non-redundant (nr) protein database that was carried out with the DIAMOND program [45]. The search results were imported into Blast2GO version 4.0 [46] for gene ontology (GO) assignments. OrthoFinder program (v2.3.3) was used to identify orthologous groups of proteins among the two de novo assemblies (MbK and Mb7) and the reference genome (GCA_013426205.1) [47].

Data availability

The raw paired-end reads were deposited in the National Center for Biotechnology Information BioProject database (http://www.ncbi.nlm.nih.gov/bioproject) with IDs PRJNA819239 and PRJNA819240 for MbK and Mb7, respectively.

Methylated-adenine library construction

Three gDNA extractions from harvested conidia were performed to construct the methylated-adenine libraries. The conidia were collected from (1) 14-d-old SDA plates with MbK, (2) 14-d-old SDA plates with Mb7, both serving as saprophytic growth samples, and (3) 3 pathogenic sample gDNA from 10 individual *Myzus persicae* aphids. Libraries were constructed as described previously [48] with some modifications. Briefly, extracted gDNA was predigested with EcoRI (Thermo Fisher Scientific) and subjected to further digestion with DpnI (Thermo Fisher Scientific). Digested gDNA served as the template for library construction using blunt-end adaptors (S1 Table, AdRt and AdRb) for PCR amplification using the adapter’s complement primers (S1 Table, AdR_PCR), then was separated on and extracted from a 1% agarose gel using the ZymoClean Gel DNA Recovery Kit (Zymo). Ligation was carried out using the CloneJET PCR Cloning Kit (Thermo Fisher Scientific), followed by transformation into DH5a competent cells (Bio-Lab, Israel). Colony PCR was conducted using the primers provided with CloneJET and positive colonies (containing an insert) were submitted to Sanger sequencing (hylabs, Israel).

Validation of swc4 gene adenine methylation

Conidial gDNA was extracted from Mb7 and MbK isolates harvested from 14-d-old SDA cultures, and evaluated by NanoDrop spectrophotometer and 1% agarose gel separation. The gDNA was diluted to 500 ng with ultrapure water (Bio-Lab) and then digested with DpnI. Untreated gDNA served as a control. The digested material served for PCR amplification using specific primers encompassing a region within the swc4 gene (S1 Table, SWC4_1 + 2 or SWC4_3 + 4). Amplicons were analysed by separation on a 1% agarose gel in the presence of ethidium bromide.

RNA extraction from fungal material and cDNA synthesis

Conidia were harvested from SDA plates in the presence of 0.01% (v/v) Triton X-100 in distilled water.
Samples were flash-frozen in liquid nitrogen with the addition of TRIzol (Thermo Fisher Scientific) and ground with metal beads in the Geno/Grinder at 6,500 oscillations/min for 3 min. Then, total RNA extraction was carried out according to the TRIzol manufacturer’s protocol. The extracted samples were treated with DNaseI (Thermo Fisher Scientific) to remove gDNA according to the manufacturer’s instructions. Total RNA was analysed by NanoDrop spectrophotometer and 500 ng were taken for cDNA synthesis with an oligo dT primer using the Verso cDNA Synthesis Kit (Thermo Fisher Scientific) according to the manufacturer’s instructions.

**Differential expression determination by RT-qPCR**

The obtained cDNA was subjected to qRT-PCR using Fast SYBR Green Master Mix (Thermo Fisher Scientific) and tef as the reference gene using specific primers ([S1 Table](#S1_Table), qMb_TEF F and R [49]). The primers for amplifying swc4 ([S1 Table](#S1_Table), qMb_SWC4 F and R; qMb_SWC4 p.c. F and R) and swr1 ([S1 Table](#S1_Table), qMb_SWR1 F and R; qMb_SWR1 p.c. F and R) were examined for their efficiency by conducting standard curve analysis and then served for qRT-PCR analysis for differential expression. The qRT-PCR analyses were conducted three times for each comparison. The relative expression levels were calculated using the $2^{-\Delta\Delta Ct}$ method [50]. The relative expression results were statistically analysed by standard least squares analysis by restricted maximum likelihood (REML), using repeat as random effect, followed by post hoc comparison by Student’s t-test using JMP Pro software version 16.0.0 (SAS Institute Inc.).

**CRISPR single guide (sg) RNA design for swc4 and swr1 gene mutagenesis, Cas9 reaction setup**

The CRISPR mutagenesis protocol was based on Davis et al. [51]. Single guide RNA targeting genes of interest was designed using Benchling [Biology Software (2021) Retrieved from https://benchling.com] based on Cas9 cutting efficiency [52]. The sgRNA template was synthesized as a primer ([S1 Table](#S1_Table), SWC4.sg132 or SWR1.sg1979) and served for sgRNA synthesis using the EnGen sgRNA Synthesis Kit (NEB, USA) following the manufacturer’s instructions. The sgRNA was then purified using the RNA Clean & Concentrator-25 Kit (Zymo) following the manufacturer’s instructions. RNA concentration and quality were assessed by NanoDrop spectrophotometer. For each CRISPR reaction: 2.5 µL concentrated sgRNA, 2 µL of EnGen Spy Cas9 NLS (NEB) in the presence of buffer 3.1 (NEB) in a final volume of 5 µL were incubated at 37°C for 30 min and kept on ice until use.

**Selective double-stranded (ds) DNA marker production**

Using pSK1019 (kindly provided by Prof. Seogchan Kang at The Pennsylvania State University, USA) as a template, the regions of the egfp reporter gene (including the promoter from the *Cochliobolus heterosporus* glyceraldehyde 3-phosphate dehydrogenase gene) and *hph* (including the promoter from *Aspergillus nidulans trpC*) were amplified using primers pSK_DS_HYG_F and pSK_DS_egFP_R ([S1 Table](#S1_Table)), designed to overlap with ca. 100 bp upstream and downstream of the region resulting in a 2946-bp amplification. For the PCR: 1 µL of pSK1019 plasmid (20 ng/µL) and 1 µL of each of the primers (10 µM) were amplified using Phusion High-Fidelity DNA Polymerase (NEB) in a final volume of 20 µL according to the manufacturer’s instructions, under the following conditions: 98°C for 30 s followed by 35 cycles of: 98°C for 10 s →60°C for 10 s →72°C for 3 min, and a final extension step at 72°C for 5 min. The PCR product was separated on a 1% agarose gel, and purified using the Zymoclean Gel DNA Recovery Kit. The concentrated product (ca. 200 ng/µL) served as the selective dsDNA marker for CRISPR mutagenesis.

**Fungal protoplast transformation and colony selection**

Protoplasts of *M. brunneum* MbK were prepared as described previously [51,53] with some modifications. Briefly, conidia harvested from a 7-d-old culture of isolate MbK served as a fungal starter, which was grown for 3 d in malt extract broth media. Then, the mycelium was filtered through miracloth (Merck Millipore, USA), washed with sterile distilled water and with KCl/CaCl$_2$ buffer (KCl 89.92 g/L, CaCl$_2$ 7.35 g/L), and transferred into a flask containing 20 mL of enzyme solution [50 mL KCl/CaCl$_2$ buffer containing 200 mg lyzing enzyme (Sigma, USA), 150 mg driselase (Sigma), 15 mg lyticase (Sigma), 10 mg bovine serum albumin (Sigma), and 10 mg yatalase (Takara Bio, Japan)] and incubated for 2–3 h at 28°C with shaking at 90 rpm/min. Protoplasts were separated from the mycelial debris by filtration through miracloth and washed twice with KCl/CaCl$_2$ buffer, then suspended in 750 µL STC buffer solution (1.2 M sorbitol, 100 mM Tris pH 7.5, 0.05 M CaCl$_2$-2 H$_2$O), and the concentration was estimated by haemocytometer. STC buffer was added to obtain a final concentration of $5 \times 10^7$
protoplasts/mL. A 1/4 volume of PEG solution [two volumes of 37% v/v PEG 8000, with one volume of 3X PEG amendments (1.8 M KCl, 150 mM CaCl₂, 150 mM Tris pH 7.4)] was added to the protoplasts in the STC buffer. For transformation, 125 L of protoplast solution was incubated with 5 L of a sgRNA – Cas9 mix and 5 L of the selective dsDNA on ice for 30 min. Then, 1 mL of PEG solution was added and the mixture was incubated for an additional 20 min at room temperature. Transformed protoplasts were plated on 10 mL growth medium (sucrose 239.4 g/L, yeast extract 0.5 g/L) and left to recover for 2–3 h, then the top layer of 20 mL media containing Hygromycin B Gold antibiotic (InvivoGen, France) at a final concentration of 500 μg/mL per plate, was added and left to dry. Plates were incubated at 28°C for 2 weeks, during which time emerged colonies were collected and transferred to SDA plates containing 200 μg/mL Hygromycin B Gold. Colonies were then single-spore isolated and served for gDNA extraction followed by PCR amplifications and sequencing of the swc4 or swr1 genes.

**Morphological characterization of mutants**

Fungal isolates Mb7 and MbK and the swc4 and swr1 mutants generated from MbK were used for morphological analysis of growth. Using a 200-μL sterile tip, a round piece of fully sporulated SDA (5 mm) was collected from 14-d-old SDA plates, placed in 5 mL 0.01% Triton X-100 and vigorously vortexed to obtain a conidial suspension. The conidial concentration was estimated by haemocytometer. Each suspension was then diluted to a final concentration of 1 × 10⁶ conidia/mL. A 30-μL aliquot of the diluted suspension was spread on an SDA plate for germination assay, which was examined 16 h postplating. Simultaneously, 30-μL aliquots were placed in the middle of another three SDA plates, without spreading, for radial growth assay. Plates were maintained as detailed in the “Fungal growth and maintenance” section and growth was documented at 7 and 14 d postplating. The assays were performed at least three independent times for each fungal isolate and mutant. Statistical analysis was performed using JMP Pro software version 16.0.0. Conidial concentration, germination and radial growth were subjected to standard least squares analysis by restricted maximum likelihood (REML) method followed by Dunnett’s test for conidial concentration and germination using MbK as control group for comparison. Tukey’s HSD test for multiple comparisons among means was used for radial growth.

**Metabolomic analysis**

**Fungal growth conditions**

Fourteen-day-old fungal SDA plates of Mb7, MbK, and MbK-swc4-mutant served as the source material for the metabolomics analysis and metabolite assay. A 1 cm × 1 cm piece from each plate was placed in a 250-mL flask containing 100 mL of sterile SDB (Difco). Inoculated flasks were sealed with sterile aluminium foil and cultured at 28 ± 0.5°C in an incubator in the dark for stationary growth for 30 d [54]. In total, five repeats of Mb7 and MbK were used, and three of MbK-swc4-mutant. At the end of incubation, liquid from each flask was separately transferred through miracloth into a sterile container, separated into two 50-mL tubes and kept at −20°C for further analysis. A single tube from each repeat was subjected to LC – MS analysis, while the other served as a pool for insect-injection bioassays.

**LC – MS analysis**

A 1-mL aliquot of each biological replicate was used for spectrometric analysis, using an Agilent HP 1260 Infinity Series liquid chromatography instrument coupled to a Q-TOF mass spectrometer with a Dual ESI source of ionization, and equipped with a DAD system (Agilent Technologies, USA) following a previously described setup [55]. An Adamas C-18 column (4.6 × 50 mm, 3.5 μm, SepaChrom, Italy) was used for the chromatography.

**Statistical analysis of LC – MS data**

Statistical analysis of the metabolomics data was carried out by Mass Profile Professional software, version 13.1.1 (Agilent Technologies). Raw data of biological and technical replicates were grouped by isolate and mutant (Mb7, MbK and MbK-swc4-mutant) and the groups were subjected to one-way ANOVA and FC ≥ 2.0 analyses. The results were then subjected to PCA and depicted as hierarchical clusters. Relevant statistically significant compounds were identified using an in-house fungal metabolite database, and by comparison with data available in the literature.

**Insect rearing and bioassay**

*Galleria mellonella* larvae were reared in sterilized glass jars, fed every 2–3 d and occasionally split to avoid overcrowding and disease development. The jars were
maintained in a 25 ± 2°C chamber, with a 12:12 h light regime and an air drier to maintain low humidity.

**Mortality bioassay in *G. mellonella***

For fungal virulence bioassay of MbK and Mb7 compared to the mutants, conidia were collected from 14-d-old SDA culture plates and mixed in sterile 0.01% Triton X-100 to obtain a uniform spore suspensions. The suspensions concentration were estimated by haemocytometer. Suspensions were then diluted to obtain 1 × 10^7 conidia/mL for inoculum. Experiments were conducted in 90-mm petri dishes, for inoculation Whatman filter paper wetted with 1 mL of spore suspension or 0.01% Triton X-100 as a negative control was placed within the plates. Five to eight *G. mellonella* 5th-stage larvae were placed in each petri dish and left to acquire conidia from the inoculated filter paper. Each treatment was conducted in five experimental plates. All plates were placed in a box with pre-wetted paper on the bottom, sealed with a plastic bag and maintained in a 25°C incubator. Mortality was assessed daily (larvae were detached from their silk and returned to the plate). Cadavers were surface-sterilized with bleach and 70% ethanol, washed with distilled water and kept for conidiogenesis in separate petri dishes in a 28°C incubator. The entire set of experiments was conducted in five independent repeats. For statistical analysis, arcsine-transformed values of mortality were subjected to standard least squares analysis by restricted maximum likelihood (REML) method followed by Tukey HSD post hoc analysis for multiple comparisons among means using JMP Pro software version 16.0.0.

**Injection of cultural filtrates into *G. mellonella***

The pooled metabolite solutions were used for injection bioassays in 5th-stage *G. mellonella* larvae to assess the toxicity of each fungal filtrate. Each larva was inoculated by injection of 10 μL suspension into the pre-last proleg and placed on a moistened Whatman filter paper in a 90-mm petri dish. Injected larvae were monitored for the first 10 min to remove paralysed individuals, which were replaced with new treated larvae. Each treatment was conducted in three technical repeats, each containing 10 injected larvae. As a negative control, larvae were injected with 10 μL sterile SDB. A mock treatment contained three repeats of 10 un-injected larvae. The plates were placed in a box, sealed in a plastic bag and incubated in a 25°C incubator. The entire set of bioassays was conducted three times. Mortality measurement and statistical analysis were performed as described above.

**Results**

**Isolates MbK and Mb7 demonstrate low genetic polymorphism**

**Resequencing analysis**

The obtained data enabled to compare MbK and Mb7 genomes and to seek for nonsynonymous mutations which might explain phenotypic differences observed previously [7]. In the analysis, a small number of single-nucleotide polymorphisms (SNPs) and base insertions/deletions (INDELS) were identified that distinguished between the isolates and the reference genome (*Metarhizium brunneum* 4556; ASM1342620v1). In total, 269 and 270 mutations were found in Mb7 and MbK compared with the reference, respectively. Among these mutations, 248 were present in both isolates including six frameshift variants, four missense variants and one synonymous variant (Figure 1(a); S1 file). The 43 variations distinguishing between the isolates were upstream or downstream of predicted genes (length of 5Kb). None of the identified variations, were located within coding sequence (CDS), and 11 mutations (5 in Mb7 and 6 in MbK) were found to be ±1Kb of predicted genes (Table 1; Figure 1(b)).

**de novo analysis**

Further, we conducted *de novo* analysis in order to identify structural differences that might be missed in the variant calling results. The clean reads were assembled into 348 and 284 scaffolds, for the genomes of Mb7 and MbK isolates, respectively. These were used for gene prediction which revealed minor differences between the isolates, 10,532 genes were identified in Mb7 genome and 10,555 in MbK genome. Comparison of orthogroups between the reference genome and the two isolates, resulted in possible duplications or deletions (S2 file). In the analysis, we detected 56 orthogroups with different copies of genes between the isolates, out of these 40 orthogroups were present only in one isolate: 16 orthogroups in MbK and 24 in Mb7 (Figure 2(a)). Blast2GO analysis on the genes within the unique orthogroups, revealed high similarity in function. Primary metabolic processes demonstrated similar percentage in both isolates (19% and 20% in Mb7 and MbK, respectively), while regulation of primary metabolic processes was identified only within MbK unique orthogroup (S1 Fig). Genes encoding Trypsin were found in two orthogroups, OG0000446 – containing only 1 gene in Mb7 isolate, and OG0000103 – containing two genes in Mb7 and 3 genes in MbK isolate (S2 File). Interestingly, the orthogroup related to protease enzymes
(OG0000096) included three genes in the reference and in MbK isolate, while only two genes were identified in Mb7 genome (Figure 2(b); S2 File). Similar protease activity was not found in other orthogroups.

Adenine methylation in M. brunneum isolates

Methylated-adenine sequence libraries were constructed from MbK and Mb7 genomic (g) DNA. Based on the obtained size variation in the colony PCR performed on the obtained bacterial colonies (S2 Fig), 102 colonies were selected for further sequencing. Of these 102 sequences, 38 were collected from an infected aphid library, which mainly provided bacterial sequences from aphids' symbionts. Other libraries provided all M. brunneum-related sequences and all, except one, were sequences within genes (a partial list of obtained genes is presented in S2 Table).

Differential methylation and expression of swr1 complex protein 4 (swc4) between fungal isolates and growth stages

Further analysis was carried out on the swc4 gene, which was identified in the saprophytic conidia library of the MbK isolate. Observation of the known sequence of swc4 revealed nine sites of GATC which might be methylated on adenine and detectable by DpnI digestion (Figure 3(a)). Amplification using SWC4_1 and SWC4_2 primers, generated a 324-bp amplicon which was demonstrated repeatedly in both digested and undigested gDNA from Mb7 and MbK isolates, meaning no cleavage by DpnI and thus no adenine methylation on the GATC site (Figure 3(b)). In the amplification using SWC4_3 and SWC4_4 primers, generating a 1248-bp amplicon, differential amplification was observed. Whereas in the Mb7 isolate, similar amplification was detect in digested and undigested gDNA samples, in all three repeats conducted on the MbK isolate, lower amplification of this region was detected in the DpnI-digested gDNA (Figure 3(c)). Digestion of the same gDNA with DpnII, which detects unmethylated GATC sites, revealed no amplification in any of the repeats (data not shown). This might have resulted from one GATC site not being methylated. To examine the expression of swc4 and to link the detected methylation to gene regulation, we conducted quantitative (q) RT-PCR on MbK and Mb7 RNA extracted from conidia. MbK demonstrated expression of swc4, however with 60% reduction compared to Mb7 (t-test: DF = 1, t Ratio = 10.25, p < 0.0001, Figure 3(d)).

Figure 1. Genetic polymorphism between Metarhizium brunneum isolate, Mb7 and MbK. (a) Venn diagram comparing identified mutations in isolates MbK and Mb7 based on reference genome Metarhizium brunneum 4556 (ASm13426201), conducted by Venny 2.1 tool [56]. (b) Representative snapshot from IVG software showing deletion in MbK genome compared with reference genome and Mb7 isolate. Location in reference genome CP058935.1 nt 101,270. gff: represents gene annotations, upstream to XP_014539572.1 (MEPI) and downstream to XP_014539573.1 (PRIC) (c) Enlargement of IGV snapshot of the deletion in MbK isolate compared with reference genome and Mb7 isolate.
Table 1. SNPs and INDELs located near predicted genes distinguishing between *M. brunneum* isolates MbK and Mb7.

| Location Chromosome (position) | Nearby gene (up- and downstream from mutation) Accession no. of predicted protein in *M. brunneum* | Distance (bp)* | Reference genome** | Mb7 | MbK |
|--------------------------------|-------------------------------------------------------------------------------------------------|----------------|--------------------|-----|-----|
| CP058932.1 (5,610,772)        | QLI66165.1 (hypothetical) XP_014543791.1 (stress response protein nst1) XP_014546251.1 (lipase A) XP_014548845.1 (secretory lipase) | +1083          | C                  | CT  | C   |
|                               | QLI65247.1 (hypothetical) XP_014541214.1 (oligopeptide transporter)                            | +943           |                    |     |     |
| CP058933.1 (7,300,466)        | QLI67091.1 (grayanic acid biosynthesis cluster O-methyltransferase) XP_014545711.1 (O-methyltransferase) QLI66338.1 (serine/threonine-protein kinase gad8) XP_014529469.1 (protein kinase-like protein) | +4,272         | C                  | CTT | C   |
| CP058934.1 (89,044)           | QLI69718.1 (pyranose dehydrogenase) XM_014684893.1 (glucose-methanol-choline oxidoreductase) QLI69015.1 (monodictyphenone cluster transcription factor) XP_014540380.1 (aflatoxin biosynthesis regulatory protein) | −119           | G                  | G   | A   |
| CP058934.1 (1,140,468)        | QLI70014.1 (alpha-ketoglutarate-dependent taurine dioxygenase) XP_014544201.1 (taurine catabolism dioxygenase tauD/TfdA) QLI69251.1 (nonribosomal peptide synthetase fmpE) XP_014544202.1 (pyridoxal phosphate-dependent transferase) | −821           | G                  | G   | C   |
| CP058934.1 (3,332,604)        | QLI70028.1 (hypothetical) XP_014542631.1 (FluG domain-containing protein)                     | +371           | C                  | T   | C   |
| CP058934.1 (3,332,605)        | QLI69264.1 (hypothetical)                                                                    | −246           | A                  | AATT| A   |
| CP058934.1 (3,332,610)        | QLI71021.1 (hypothetical) XP_014539572.1 (MEP1)                                             | −861           | AGCGGACGTC         | AGCGGACGTC | A   |
| CP058935.1 (101,270)          | QLI71289.1 (minor extracellular protease vpr) XP_014539573.1 (PR1C)                          | +1150          |                    |     |     |
| CP058936.1 (4,024,430)        | QLI71985.1 (hypothetical) XP_014541896.1 (ankyrin repeat protein)                            | −1023          | T                  | T   | TGGGGG GGGGGGGGG |
| CP058937.1 (280,063)          | QLI73557.1 (hypothetical) XP_014540789.1 (hypothetical) QLI73412.1 (hypothetical)          | +522           | C                  | C   | T   |

*Location of mutation in base pair upstream (−) or downstream (+) of gene.

**Reference genome: *Metarhizium brunneum* 4556 (ASM1342620v1).
Cas9-mediated transformation and mutant selection

Mutagenesis of swc4 was performed with guided RNA located at nt position 132 (S1 Table, SWC4_sg132) and repeated four independent times. Out of over 100 mutants collected, 26 were further transferred to new Sabouraud dextrose agar (SDA) plates containing the antibiotic hygromycin. Eight colonies were unable to grow and were discarded. The other 18 colonies were used for gDNA extraction and sequencing of the swc4 region in the genome. Only one single colony did not demonstrate integration of the construct hygromycin phosphotransferase-green fluorescent protein gene (hph-gfp) within swc4. All other mutants had a similar...
integration site (at 129 nt in swc4) with similar distribution of insert orientations (10 in the swc4 orientation and 7 in reverse orientation). Some of the mutants demonstrated GFP emission under a confocal microscope. However, the signal was too low to be useful as a marker (data not shown). Most of the clones had a deletion in the gfp gene. In all mutants, the gene that was oriented as swc4 (either gfp or hph) had a deleted stop codon. None of the examined mutants were able to cause a frameshift in swc4 or to add an early stop codon; rather, the reading frame was left intact after the integration site and until the native stop codon of swc4 (Figure 4(a)). A single repeat was carried out using another guided RNA at nt position 538 which is located in the SANT domain of the protein. None of the examined colonies demonstrated integration of the insert in swc4, and this reaction was therefore not repeated (data not shown).

Mutagenesis of the non-essential SWR1 protein, which is part of the same cellular complex [57], was carried out with guided RNA designed for nt position 1979 in swr1 (S1 Table, SWR1_sg1979). Twenty-one colonies were collected following transformation, of which a single colony was unable to grow on SDA amended with hygromycin. Two colonies were further used for sequencing, and one was confirmed to have the insert integrated at nt position 1979 in swr1 (Figure 4(b)), whereas in the other colony, no integration into swr1 was detected. The selected positive colony had no deletions in gfp, hph or swr1. The stop codon of hph was demonstrated in the sequence. Insertion of 63 and 78 bp was detected surrounding the insert downstream of the stop codon of gfp and hph, respectively. GFP signal was undetectable by confocal microscopy (data not shown).

Analysis of swc4 expression in the mutant (downstream of the insertion, S1 Table, qMb_SWC4 F + R) compared to isolates Mb7 and MbK revealed down-regulation, although it was not significant. In the swr1 mutant (downstream of the insertion, S1 Table, qMb_SWR1 F + R), the gene was significantly downregulated (F[2,4] = 22.86, p = 0.0065; Tukey HSD p < 0.05) (S3 Fig). In parallel, amplification with primers encompassing the insertion site in both swc4 and swr1 (S1 Table, qMb_SWC4 p.c F + R and qMb_SWR1 p.c F + R) occurred only in the isolates Mb7 and MbK (in swc4 gene: F[2,4] = 1242.32, p < 0.0001, t-test for all comparisons p < 0.0001; in swr1 gene: F[2,4] = 30.73, p = 0.0059, t-test comparison: Mb7 vs swr1-mutant p = 0.0029 and MbK vs swr1-mutant p = 0.0069) (S3 Fig).

Reduced fungal saprophytic growth in swc4 and swr1 mutants

Morphological characterization of the mutants included visual observation of conidial morphology and sporulation rate, germination and radial growth estimates. During the growth of both fungal mutants, none of the vital processes were arrested; all fungi

**Figure 4.** Schematic representation of the mutants generated in this study. (a) swr1 complex protein 4 (swc4) with integration of hph-gfp starting from nt position 132 of swc4, hph stop codon deleted, and swc4 remaining in-frame until the native stop codon at nt position 1972 of the gene. (b) swr1 gene with integration of hph-gfp starting from nt position 1978, hph stop codon present at its original location and before nt position 1979 of swr1. Top yellow-shaded boxes in both panels represent PCR sequence alignments.
completed cycles of sporulation, germination and hyphal elongation, but with a significant impact on each stage. Conidial production was significantly reduced by one order of magnitude in the MbK-swrl-mutant and MbK-swrl-mutant compared to MbK and Mb7, while no difference observed between Mb7 and MbK (F(3,16.08) = 12.05, p = 0.0002; Dunnett’s test: swrl-mutant p = 0.0003 and swrl-mutant p = 0.003) (Figure 5(a)); however, conidial size and shape did not change (Figure 5(b)). Conidial germination was significantly reduced in both mutants compared to Mb7 and MbK isolates which displayed no difference in germination (F[3,17] = 11.86 p = 0.0002; Dunnett’s test: swrl-mutant p=0.032, swrl-mutant p = 0.0001) (Figure 5(c)). Hyphal elongation and branching were delayed in both mutants compared to Mb7 and MbK (F (3,26.34) = 520.38, p<0.0001; Figure 5(d)) and at both 7 and 14 d of growth, radial colony size was reduced in the swrl mutant compared to Mb7 and MbK (Tukey HSD: p < 0.0001). The swrl mutant demonstrated the most delayed growth, and the difference was significant compared to MbK, Mb7 and the swrl mutant (Tukey HSD: p < 0.0001) (Figure 5(e,f)).

**Reduced virulence of swrl and swrl mutants in Galleria mellonella**

To assess the impact of the swrl and swrl genes on virulence and pathogenicity, bioassays were conducted with the mutants on G. mellonella 5th stage larvae. The two mutants were compared to the MbK and the Mb7 isolates. The fungal source material was at 10^7 spores/mL and germination rates were above 85% in all treatments and repeats. MbK was already able to cause significant mortality of 15% at 3 d postinoculation (DPI; F(4,107.4) = 5.16, p = 0.0008). Mb7, MbK-swrl-mutant and MbK-swrl-mutant caused 19%, 24%, and 8% mortality at 5 DPI, respectively, while the MbK isolate caused over 50% mortality at this time point (F (4, 98.03) = 24, p < 0.0001). Finally, MbK caused the highest mortality rate at 7 DPI (90% on average, F (4,88.19) = 70.59, p < 0.0001), whereas Mb7 and MbK-swrl-mutant caused 75% and 73% mortality, respectively. Mortality of only 27% was achieved with the swrl mutant and the difference was not significant compared to the negative control (Fit least square: between isolates (treatment, F(4,739.4) = 75.63 p < 0.0001), during time (DPI, F(1,373.9) = 593.69 p <

**Figure 5.** Characterization of fungal morphology and growth. (a) Conidial concentration (1 × 10^6 conidia/ml ± SE) of the fungal isolates, in a 5-mm agar piece/5 mL Triton X-100 as counted by haemocytometer. (b) Conidia visualized in haemocytometer (enlarged photo of light microscopy, ×400 magnification). (c) Germination rate (% ± SE) of 1 × 10^6 conidia/ml 16 h after spreading on an SDA plate. (d) Germinated conidia, 16 h after spreading of 1 × 10^6 conidia/ml on an SDA plate (visualized using light microscopy at ×400 magnification, bar size: 50 µm). (e) Radial growth (cm ± SE) after placing 1 × 10^6 conidia/ml on the centre of an SDA plate. (f) Radial growth on SDA plates (60 mm) at 7 d (top images) and 14 d (bottom images). Statistical analysis was carried out by Tukey – Kramer test, significant differences are marked with different letters (p < 0.05).
0.0001) and within the treatment*DPI interaction (F (4,736.9) = 67.84, p < 0.0001). (Figure 6).

**Metabolomic differences between MbK, Mb7, and MbK-swc4-mutant**

To further evaluate differences that might explain the variations in virulence, we analysed the metabolomes of the two isolates (Mb7 and MbK) and the swc4 mutant. Filtrates from Mb7, MbK and MbK-swc4-mutant cultures were subjected to LC – MS analysis. Using the resulting data, including total ion chromatograms (Figure 7(a)) and mass spectra, and by comparison with an in-house fungal database and reported data [58], SMs produced by the *Metarhizium* spp. were identified (Table 2). The data included swainsonine, serinocyclin A and B, and a set of destruxins [59,60]. Statistical analysis of the complete SM data from each of the fungal isolates enabled their separation into three groups by principal component analysis (PCA) (Figure 7(b)). In this analysis, 95% of the differences (PC1) contributed to a major separation between MbK and both Mb7 and MbK-swc4-mutant. The difference observed between Mb7 and MbK-swc4-mutant contributed 5% (PC2, Figure 7(b,c)). In total, the levels of 538 SMs were detected as significantly different between at least two samples (S3 File). Fold-change (FC) analysis between the samples revealed: (a) 503 (93%) metabolites with common levels between Mb7 and Mbk-swc4-mutant (−3 < log FC < 3); (b) 521 (97%) SMs that were at higher levels in MbK compared to Mbk-swc4-mutant and (c) 506 (94%) SMs that were at higher levels in MbK compared to Mb7 (S3 File). However, among all significantly different metabolites, three were identified by comparison with a fungal database and with the literature (Table 3).

**Pathogenic activity of Mb7, MbK, and MbK-swc4-mutant culture filtrates**

To evaluate the pathogenic potential of the different filtrates containing SMs from Mb7, MbK, and the mutant of swc4, we exposed *G. mellonella* larvae by injecting sterile filtrate directly into their haemocoel without the presence of fungal conidia. A significant difference was observed as early as 3 DPI: larvae injected with MbK filtrate were consistently dead whereas no mortality was observed with the other treatments (F(4,32.58) = 371.88, p < 0.0001). Severe melanization occurred on larvae injected with MbK filtrate and was already detectable 1 DPI. At 5 DPI, melanization was also noticeable in some individuals in the other two treatments, but their motility was not affected (Fit least squares: treatment F(4,158.5) = 110.5 p<0.0001, DPI F(1,158.6) = 177.25 p < 0.0001, treatment*DPI F(4,158.5) = 102.99 p < 0.0001) (Figure 8).

**Discussion**

We previously showed intra-species variation in disease progression and pathogenicity against aphids between *M. brunneum* isolates MbK and Mb7 [7]. Here, we compared the genetic and epigenetic variation among the isolates to elucidate their role in the varied virulence.

![Figure 6](image-url) Cumulative mortality (% ± SE) of *G. mellonella* inoculated with spore suspensions of *M. brunneum* isolates MbK, Mb7 and mutants of the genes swr1 and swc4 for 7 d postinoculation. Statistical analysis were carried out on arcsine square root-transformed values of mortality proportion using Tukey – Kramer test; significant differences are indicated with different letters (p <0.05). On the right: representative larvae infected with (from top to bottom): MbK, Mb7, MbK-swc4-mutant, MbK-swr1-mutant and non-infected larvae.
When the genomes of both isolates were compared, we found unexpectedly low genetic variation (SNPs, INDELs) that was evenly dispersed throughout both genomes; we did not detect any highly polymorphic regions [61]. A much higher percentage of polymorphisms between fungal isolates has been observed by others, accompanied by nonsynonymous mutations [61,62]. All of the mutations differentiated MbK from Mb7 were located in intergenic regions. However, mutations located near predicted genes might have a cis-regulatory role [63]. An interesting candidate for this type of gene-expression regulation, which might have a role in pathogenicity, is the identified mutation located between two genes, pr1c and mep1, encoding proteases. These genes play a key role in *Metarhizium*’s pathogenic lifestyle acting as virulence factors [64,65,66]. Moreover, differential expression was observed in mep1 gene in response to antifungal agents [6]. Furthermore, de novo assembly of Mb7 and MbK genomes revealed high similarity in gene number and functional gene groups. Mainly, absence of specific genes in one of the isolates was a result of grouping similar gene into different orthogroups rather than gene deletion. Interestingly, pathogenicity-related

![Figure 7. Metabolomic comparison between Mb7, MbK and MbK-swC4-mutant cultured in Sabouraud dextrose broth (SDB). (A) Total ion chromatograms of the culture filtrates from Mb7 (top), MbK-swC4-mutant (middle) and MbK (bottom). Data were recorded in positive ionization mode. (B) PCA score plot of the LC – MS data acquired in positive mode. Mb7 replicates are depicted in blue; MbK replicates in brown; and MbK-swC4-mutant replicates in red. (C) Hierarchal condition tree heatmap of differential metabolic profiles from *M. brunneum* culture filtrates. The abundance of each compound is associated with a colour ranging from blue (less abundant) to red (more abundant).](image)
protease orthogroup demonstrated deletion of one gene in the Mb7 isolate. This deletion have to be further studied, possibly by genome editing, in order to elucidate its role in the varied virulence between Mb7 and MbK isolates.

In an attempt to identify an epigenetic variation, we detected 6 mA in the *M. brunneum* genome of both isolates. Specifically, we found the differential presence of 6 mA on swc4 gene, which was correlated with reduced expression in MbK. We hypothesized that 6 mA plays a regulatory role in swc4, which in turn can affect a cascade of events by chromatin remodelling of regions regulating pathogenicity. *Metarhizium* growth as a pathogen requires an arsenal of enzymes and metabolites to conquer the host, especially as the host utilizes rapid epigenetic changes it order to achieve resistance [19]. The fungi should provide a flexible and rapid response which similarly may be achieved through epigenetic modifications [67]. However, in recent publications, the level of 6 mA in true fungi has been shown to be extremely low and its epigenetic regulatory role has been questioned [23,68]. The presence of 6 mA in MbK but not in Mb7 correlated with reduced expression of swc4 in MbK, as opposed to previous studies [23,69]. This might have been due to other regulatory factors known to work simultaneously [69]. The high distribution of 6 mA in the genomes of prokaryotes, early-diverging fungi, ciliates and algae.

**Table 2.** Identified metabolites of *Metarhizium* spp. obtained by LC – MS analysis.

| Compound        | RT (min)* | Experimental mass (Da) | Theoretical mass (Da) | Molecular formula |
|-----------------|-----------|------------------------|-----------------------|-------------------|
| Swainsonine     | 1.033     | 173.106                | 173.106               | C17H11N06         |
| Serinocyclin A  | 1.356     | 672.3094               | 672.3094              | C32H44N12         |
| Serinocyclin B  | 1.477     | 656.3179               | 656.3179              | C32H44N11         |
| Destruxin E-diol| 4.348     | 611.3531               | 611.3531              | C32H44N10         |
| Destruxin D     | 5.537     | 623.3525               | 623.3525              | C32H44N10         |
| Destruxin E     | 5.605     | 593.3432               | 593.3432              | C32H44N8          |
| Destruxin C2    | 6.129     | 595.3524               | 595.3524              | C32H44N8          |
| Destruxin A     | 6.134     | 577.3454               | 577.3454              | C32H44N7          |
| Destruxin B2    | 6.406     | 579.3588               | 579.3588              | C32H44N7          |

*Retention time.
†Mono-isotopic mass.

**Table 3.** Significant differences in metabolites between filtrates from Mb7, MbK, and MbK-swcd4-mutant (swc4) cultured in SDB.

| Compound         | (swcd4 vs. Mb7) | (swcd4 vs. MbK) | (Mb7 vs. MbK) | Differential |
|------------------|-----------------|-----------------|---------------|--------------|
| Swainsonine      | 0.84            | -16.83          | -17.68        | Mb7, swc4 < MbK |
| Serinocyclin A   | 17.97           | 0.39            | -17.58        | Mb7 < MbK, swc4 |
| Serinocyclin B   | 17.10           | -0.45           | -17.56        | Mb7 < MbK, swc4 |

**Figure 8.** Cumulative mortality (% ± SE) of *G. mellonella* inoculated with filtrate collected from *M. brunneum* isolates MbK, Mb7 and MbK-swcd4-mutant during 5 d postinjection. Statistical analysis was carried out on arcsine square root-transformed values of mortality proportion using Tukey – Kramer test, and significant differences are indicated by different letters (p < 0.05). On the right: representative larvae injected with (from top to bottom) MbK, Mb7 and MbK-swcd4-mutant filtrate. SDB, Sabouraud dextrose broth.
enables a detailed study of its role in regulating gene expression [23,70,71]. Conversely, in animals, plants and true fungi, further study is required to understand the mode of action and role of 6 mA, despite its low presence in their genomes [68].

Notably, *swc4* might be downregulated to some extent in the MbK isolate and may provide better activity of virulence genes and metabolite secretion, but damaged activity will result in the loss of these abilities as demonstrated in the MbK-*swc4*-mutant. The role of SWC4 subunit, as well as of SWR1-C as a regulator of biological processes in fungi is still largely unknown [32]. Complete knockout of *swc4* was futile, as it repeatedly failed using Agrobacterium tumefaciens-mediated transformation by homologous recombination (data not shown). Attempts to mutagenize its locus by Cas9 resulted in chimeric RNA of *swc4* and *hph* lacking the latter’s stop codon. Putatively, this results in fused protein which might affect SWC4 activity. In contrast, similar mutagenesis of the non-essential *swr1* gene resulted in the chimeric RNA but without the stop codon deletion that repeatedly occurred in *swc4* mutagenesis. These results support the previously reported essential nature of *swc4* [72,73,74,75]. The mutagenesis conducted in this study resulted in significant and extensive damage to the fungus. It may be that the phenotype observed in the *swc4* mutant is due to SWC4 subunit difficulty to bind to chromatin, however that requires further research in the protein level. Interestingly, this type of Cas9-mediated transformation can be a useful tool in the study of essential genes which otherwise cannot be manipulated. Moreover, this method provides a bypass for the homologous recombination limit in fungal species possessing Ku proteins for non-homologous end-joining DNA repair [76]. Thus, we present a new venue for studying essential genes.

The results of this study suggest that *swc4* has a significant role in fungal growth, virulence and metabolism, from the immediate growth attenuation in the mutant compared to Mb7 and MbK, to decreased virulence and the significant effect on SM production. It has already been proposed and sporadically proven that SMs are indeed regulated by chromatin remodelling, especially as SM genes are mainly located in clusters which are more efficiently controlled by histone rather than DNA modifications [32,34,77]. The putative disruption of SWC4 protein, and to a larger extent of SWR1-C, can shed light on its regulatory role during the fungus’ life span. As hypothesized by Chen and Ponts (2020), SWC4 mediated histone variant deposition (of H2A.Z) can play a crucial role in biological processes and secondary metabolism in filamentous fungi [32]. Indeed, the SM analysis revealed that the MbK-*swc4*-mutant produces less metabolites but interestingly, this mutation made the MbK isolate, with a single gene disruption, similar to the Mb7 isolate in terms of SM production and virulence. Allegedly, MbK isolate utilizes in parallel both toxin and growth virulence strategies; while Mb7 mainly utilize the growth strategy [7,59]. Further to this hypothesis, *swc4* disruption shifted MbK towards the growth rather than toxin strategy thus pathogenicity was attenuated to some extent but was not lost. Simultaneous elevation of growth and toxicity in a single isolate, may result in a highly virulent EPF.

As subunit SWC4 is part of both SWR and NuA4 complexes [57], whether SM regulation is based on H2A.Z deposition (SWR1-C) or histone acetylation (NuA4 complex), can only be assumed. A much more significant effect on fungal growth, morphology and virulence was observed in the non-lethal *swr1* mutant. Similar morphological effect was already observed in Candida albicans and proved to be related to the merge and separation of the two chromatin remodelling complexes, SWR and NuA4, both possessing also the SWC4 subunit [78]. The more severe phenotype in *swr1* mutant could be due to the successful integration of *hph* into the *swr1* gene resulted in RNA sequences encoding probably a truncated protein. We would expect to see, in both mutants, a similar regulatory effect on genes which are regulated by SWR1-C and not by NuA4 (as only SWC4 but not SWR1 is a shared subunit), but this needs to be studied in the future by comparing the transcriptomes of the mutants and MbK. Finally, there remains a large gap in our understanding of the role of 6 mA in gene regulation and its interaction with the SWR-C.

In conclusion, the results of the present study suggest that 6 mA mark in the essential gene *swc4* correlates with downregulation of its expression. SWC4 subunit plays a role in regulation of SMs and SWR1 subunit from the same chromatin remodelling complex is essential for virulence of *M. brunneum*. The main impact of this study lies in promoting the necessary knowledge to manipulate regulators of disease progression in order to increase the efficacy of EPF against agricultural pests. This study opens a new venue of future research possibilities, and provides a useful tool for studying one of the most interesting epigenetic regulation complexes, SWR1-C. A better understanding of SWR1-C can provide new knowledge in other fungi and organisms. This will also expand the possibilities of using genome editing to enhance the performance of *M. brunneum* and other EPF with respect to SMs secretion and pathogenicity against agriculturally important pests.
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