Effect of $H2A.Z$ deletion is rescued by compensatory mutations in *Fusarium graminearum*

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Abstract

Fusarium head blight is a destructive disease of grains resulting in reduced yields and contamination of grains with mycotoxins worldwide; *Fusarium graminearum* is its major causal agent. Chromatin structure changes play key roles in regulating mycotoxin biosynthesis in filamentous fungi. Using a split-marker approach in three *F. graminearum* strains INRA156, INRA349 and INRA812 (PH-1), we knocked out the gene encoding $H2A.Z$, a ubiquitous histone variant reported to be involved in a diverse range of biological processes in yeast, plants and animals, but rarely studied in filamentous fungi. All $\Delta H2A.Z$ mutants exhibit defects in development including radial growth, sporulation, germination and sexual reproduction, but with varying degrees of severity between them. Heterogeneity of osmotic and oxidative stress response as well as mycotoxin production was observed in $\Delta H2A.Z$ strains. Adding-back wild-type $H2A.Z$ in INRA349 $\Delta H2A.Z$ could not rescue the phenotypes. Whole genome sequencing revealed that, although $H2A.Z$ has been removed from the genome and the deletion cassette is inserted at $H2A.Z$ locus only, mutations occur at other loci in each mutant regardless of the genetic background. Genes affected by these mutations encode proteins involved in chromatin remodeling, such as the helicase Swr1p or an essential subunit of the histone deacetylase Rpd3S, and one protein of unknown function. These observations suggest that $H2A.Z$ and the genes affected by such mutations are part of the same genetic interaction network. Our results underline the genetic plasticity of *F. graminearum* facing detrimental gene perturbation. These findings suggest that intergenic suppressions rescue deleterious phenotypes in $\Delta H2A.Z$ strains, and that $H2A.Z$ may be essential in *F. graminearum*. This assumption is further supported by the fact that $H2A.Z$ deletion failed in another *Fusarium* spp., i.e., the rice pathogen *Fusarium fujikuroi*.

Author summary

The function of histone variant H2A.Z has been widely studied in *Saccharomyces cerevisiae* and higher eukaryotes. However, there is a significant lack of knowledge of the role of...
H2A.Z and its regulation in filamentous fungi. Here we demonstrate for the first time that deleterious consequences of complete removal of H2A.Z from the genome of *F. graminearum* are mitigated by genetic suppression. The suppressors detected open new possibilities to investigate in more depth the network linking H2A.Z and other genes in *F. graminearum* and used to decipher unresolved questions concerning H2A.Z.

**Introduction**

*Fusarium graminearum*, a homothallic filamentous fungus, is the major causal agent responsible for the devastating disease Fusarium head blight (FHB) of wheat, barley and other small grain cereal crops worldwide [1,2]. In wheat, FHB affects the head with symptoms of wilted kernels, reducing yield at harvest and causing billions of dollars losses [3,4]. As an additional serious concern, the presence of *F. graminearum* in kernels results in the contamination of cereals with mycotoxins, especially the extremely stable type B trichothecenes (TCTB) including deoxynivalenol (DON) and its acetylated C-15 derivatives (15-ADON). The presence of these mycotoxins on cereals, persistent in derived food and feed products, threatens the health of humans and animals [5–7]. A growing body of evidence indicates that chromatin structure changes play a critical role in the regulation of mycotoxin biosynthesis in filamentous fungi.

In eukaryotes, gene expression depends on chromatin structure. Chromatin is the fundamental packaging form of DNA, which is wrapped around an octamer of canonical core histones H2A, H2B, H3 and H4 to form nucleosomes [8,9]. Canonical histones are conserved across eukaryotic species and represent the major part of histones within an organism. The N-terminal tails of histones are extensively marked by covalent post-translational modifications (PTMs) including methylation, phosphorylation, or acetylation for example. The combinatorial positioning of these histone marks impact gene expression by altering chromatin state according to a not yet deciphered “Histone Code” [10]. Additionally, non-allelic isoforms of canonical histones called histone variants also exist in all eukaryotes [11,12]. The H2A family encompasses the largest number of variants [13]; H2A.Z is considered as the most evolutionarily conserved one [14,15]. Generally, H2A.Z and H2A have ~60% sequence similarity. It differs from H2A in the increased acidic patch and the carboxy-terminal α-helix included in the docking domain, which is a structure involved in the interaction of the H2A-H2B dimer with the (H3-H4)_2 tetramer [16,17]. H2A.Z has been identified in various species, including *Arabidopsis thaliana* [18], *Saccharomyces cerevisiae* [19], *Drosophila melanogaster* [20], and human [21]. It has been found involved in a diverse range of biological processes, including genome stability [22–24], DNA repair [25–28] and transcriptional regulation [29–31]. The absence of H2A.Z may be lethal in many organisms such as mouse [32], Drosophila [20], frogs [33] and Tetrahymena [34], but not in *S. cerevisiae* [35,36]. The dynamic process of H2A.Z deposition/removal from nucleosomes is mediated by ATP-dependent chromatin remodeling complexes, especially complexes belonging to the SNF2 superfamily [37,38]. In *S. cerevisiae*, the complex SWR1 is involved in the recruitment of H2A.Z. It can replace the H2A-H2B by H2A.Z-H2B dimers. By contrast, the removal of H2A.Z from nucleosomes is mediated by the SWR1-related Inositol requiring 80 (INO80) complex [28,39,40]. INO80 shares several subunits with SWR1 complex and its functions are usually associated with DNA double-strand breaks repair [41,42]. Homologs for H2A.Z and most subunits of yeast SWR1 and INO80 complex can be found in *F. graminearum*, meaning that similar biological processes may also occur [43].

Various studies illustrated that functions of H2A.Z are based on its collaboration with other histone marks, particularly those that decorate histone H3 tails. For example, both in human
and mouse ES cells, H2A.Z is enriched at enhancers and promoters marked by H3K4me3, a mark that activates gene expression [44,45]. Moreover, it seems that H2A.Z can act as a functional substitute for H3K9me3 in chromatin, recruiting Heterochromatin Protein 1 (HP1) when H3K9me3 levels are low for example [46]. Still in mouse ES cells, H2A.Z strongly co-localizes with H3K27me3 and Polycomb Repressive Complexes (PRCs) 1 and 2 near the transcription start site (TSS) of genes involved in cell differentiation, and helps to keep these genes silenced [47]. In filamentous fungi, major roles of chromatin structure changes by means of histone marks in regulating the biosynthesis of mycotoxins and other secondary metabolites have been evidenced. For example, in *F. graminearum*, H3K4me3 deposited by FgSet1 is required for the active transcription of genes involved in the biosynthesis of both DON and the pigment aurofusarin [48]. Consistently, lack of the COMPASS component FgCcl1, required for full H3K4me3, also results in reduced DON levels whereas aurofusarin levels remained unaffected [49]. Notably, removal of the H3K4me3-specific histone demethylase FgKdm5 also resulted in decreased DON levels, suggesting more complex regulatory circuits than previously anticipated [50]. By contrast, the histone mark H3K27me3 represses 14% of *F. graminearum*’s genome, including genes involved in secondary metabolism [51]. However, regarding the function of H2A.Z in fungi, very few studies were carried out in *Neurospora crassa* studying specifically the function of H2A.Z and identifying a role in oxidative stress response [52,53] as well as a link with the deposition of H3K27me3 [54].

Here, we hypothesized that H2A.Z plays important roles in controlling development and metabolism in two *Fusarium* spp., i.e., *F. graminearum* and *F. fujikuroi*. Using a reverse genetics approach on three different strains of each species, we provide pieces of evidence that H2A.Z may be essential in both species. Not a single homokaryotic ΔH2A.Z strain was recovered from *F. fujikuroi* wild-type strains, whereas mutants in which H2A.Z has been totally removed from the genome were obtained in all three *F. graminearum* strains. However, compensatory mutations occurred at other sites in FgΔH2A.Z mutants of different genetic backgrounds, revealing an unsuspected genome plasticity. As a whole, our results suggest a flexible network of genes that, by interacting with H2A.Z within the same functional network, can re-wire itself to rescue deleterious phenotypes.

**Results and discussion**

**Identification of FgH2A.Z as a single H2A variant in *F. graminearum***

We scanned the entire proteome of *F. graminearum* PH-1 for the presence of proteins carrying a domain characteristic of the C-terminal end of histone H2A, and found the accessions FGRAMPH1_01T26109 and FGRAMPH1_01T03973 as candidates for histone H2A and one variant. We putatively assigned FGRAMPH1_01T26109 as H2A and FGRAMPH1_01T03973 as H2A.Z by protein sequence similarity (see blastp [55] reports in S1 File and S2 File). FgH2A and FgH2A.Z are found on chromosomes 1 and 4, respectively. Gene models supported by transcriptomics data [56–58] show that both genes possess two introns, the first one being much larger in FgH2A.Z than FgH2A for which the second exon is larger (S1A Fig). As a point of comparison, the other sordariomycete *Neurospora crassa* was shown to also possess two introns in its H2A gene [59] whereas the eurotiomycete *Aspergillus nidulans* was shown to possess three ones [60]. Similarly, H2A.Z also possesses two introns in these two last species. We further examined available gene expression data obtained for asexual spores and actively growing *F. graminearum* mycelia [58]. Important changes in expression levels can be observed in FgH2A (more than five-fold, S1B Fig) but not for FgH2A.Z suggesting that, similarly to other eukaryotes [61], the expression of H2A is restricted to actively dividing cells. Finally, protein alignment found ~65% sequence identity between FgH2A and FgH2A.Z (S1C Fig), with
notably the typical H2A.Z features of an extended acidic patch and the substitution of a glutamine for a glycine at position 104 responsible for a less stable interaction between H2A.Z with H3 than H2A with H3 [17]. All together, these elements indicate that FGRAMPH1_01T03973 is a bona fide histone H2A.Z in F. graminearum.

Knock-out mutant of FgH2A.Z is viable with severe defects in development, toxin production, and stress response

To explore H2A.Z functions in F. graminearum, we proceeded with gene knock out experiments to remove the full coding sequence of FGRAMPH1_01T03973 (see Materials and Methods, and S2 Fig) from F. graminearum strain INRA349 known to be highly virulent on wheat and produce deoxynivalenol and its 15-acetylated form [62,63]. We successfully obtained one mutant I349ΔH2A.Z (verified by both PCR and southern blot, S3 Fig), suggesting that the deletion of H2A.Z in F. graminearum is non-lethal. This result is consistent with previous experiments in fungi that found H2A.Z as non-essential in N. crassa [64], S. cerevisiae [36], or Schizosaccharomyces pombe [65] for example, although they exhibited reduced growth in the deletion mutants. Our F. graminearum ΔH2A.Z mutant was indeed very slow-growing compared to its respective wild-type parental strain (Fig 1A). Strikingly, the add-back mutant I349ΔH2A.Z::H2A.Z, restoring H2A.Z expression levels to those observed in wild-type strain (see Materials and Methods, S4A Fig and S4C Fig), did not exhibit a rescued phenotype. This observation indicates that, although the single copy of H2A.Z was successfully removed from the genome, the observed effects on radial growth could not be (at least fully) attributed to the loss of H2A.Z. We also engineered a version of INRA349 in which we inserted a copy of FgH2A.Z under the control of the constitutive promoter pGPD (see Materials and Methods, and S4B Fig) and over-expressing FgH2A.Z by 7.6-fold (S4C Fig), here referred to as I349-OE::H2A.Z. No effect of radial growth could be observed in our conditions (Fig 1A).

The same experiment was repeated inoculating culture plates with 100 spores rather than a plug from a precedent culture. Similarly to our previous observation using plugs, I349-OE::H2A.Z showed similar growth to that of wild type, and growth defect in I349ΔH2A.Z compared to wild type was drastic (Fig 1B). However, the add-back mutant I349ΔH2A.Z::H2A.Z seemed to perform better and showed better ability to grow than I349ΔH2A.Z, although wild-type phenotype was not fully restored. We thus hypothesized that germination may be affected differently in I349ΔH2A.Z and I349ΔH2A.Z::H2A.Z. The kinetics of spore germination was monitored for up to eight hours in INRA349 wild type, I349ΔH2A.Z, I349ΔH2A.Z::H2A.Z, I349-OE::H2A.Z. Results are shown Fig 1C. In INRA349 wild type, germination was complete after eight hours, an observation consistent with a previously published assay using the same strain [63]. Similar results were obtained for I349-OE::H2A.Z, with 90% (± 2) of the spores germinated after eight hours. However, within the same time span, less than 20% of the spores of I349ΔH2A.Z (17% ± 4) and 42% (± 10) I349ΔH2A.Z::H2A.Z had germinated, values significantly lower than those measured for both INRA 349 wild type and I349-OE::H2A.Z. Germination in I349ΔH2A.Z::H2A.Z was higher than the one measured for I349ΔH2A.Z, although not significant (certainly due to high deviation in measurements). Considering this trend, we hypothesize that the differences in radial growth obtained above are caused by a better ability of I349ΔH2A.Z::H2A.Z to germinate and actually start growing.

We further tested the ability of our mutant strains to produce asexual spores (macroconidia). Kinetics of conidia production in a spore-inducing medium was followed for up to eight days for I349 wild-type, I349ΔH2A.Z, I349ΔH2A.Z::H2A.Z, and I349-OE::H2A.Z strains. Results are shown Fig 1D. Deletion of H2A.Z severely impeded sporulation, but phenotype was rescued in the add-back mutant I349ΔH2A.Z::H2A.Z to levels not differing significantly
from those measured in I349 wild-type and I349-OE::H2A.Z strains. Here, while I349ΔH2A.Z::H2A.Z had its ability to produce asexual spores restored, its ability to initiate germination is altered.

We pursued our exploration and investigated the ability to respond to osmotic/ionic stress and oxidative stress, which are common and major stresses for fungi [66]. We thus cultured for up to six days our INRA349 wild-type and mutant strains on CM agar plates supplemented with NaCl 1M, KCl 1M, H$_2$O$_2$ 5 mM, or H$_2$O$_2$ 15 mM, or not supplemented. Results are shown Fig 2 and S5 Fig. In terms of radial growth, wild-type strain showed little sensitivity to KCl but increased with NaCl (Fig 2A and 2B, S5A Fig, S5B Fig), although not significant over the whole growth curve (Fig 2C). Similarly, H$_2$O$_2$ had important measurable effects when applied at 15 mM rather than 5 mM (Fig 2A, 2B and 2C). Nonetheless, both NaCl and KCl caused visible discoloration of the mycelium and reduction in aerial mycelium (Fig 2B). Similiar effects were observed for I349-OE::H2A.Z, although the effects of NaCl and H$_2$O$_2$ 5 mM were this time significant (up to 6 dpi where growth curves meet, S5A Fig, S5B Fig and S5C Fig). Regarding I349ΔH2A.Z, growth curves showed different trends (Fig 2D). Treatment with NaCl and KCl both have drastic effects on growth, whereas H$_2$O$_2$ 5 mM or 15 mM showed no or moderate effects respectively. The situation is exacerbated in the add-back mutant

![Image of INRA349 mutants](https://doi.org/10.1371/journal.pgen.1009125.g001)

Fig 1. Developmental defects in INRA349 mutants. (A) and (B). Radial growth in FgINRA34 wild-type, ΔH2A.Z, ΔH2A.Z::H2A.Z, and OE:H2A.Z grown from a central 3 mm-diameter plug (A) or 100 spores (B) on CM agar for three and four days at 25°C in the dark. WT = wild-type I349; Δ = deletion mutant I349ΔH2A.Z; +H2A.Z = I349ΔH2A.Z::H2A.Z mutant; OE = I349OE::H2A.Z; dpi = days post-inoculation. (C). Germination rates of FgINRA349 wild-type, ΔH2A.Z, ΔH2A.Z::H2A.Z, and OE:H2A.Z after eight hours of incubation. (D). Fitted kinetics of sporulation followed for eight days. Black = wild-type I349; dashed red = I349ΔH2A.Z; dashed blue = I349ΔH2A.Z::H2A.Z mutant; dashed purple = I349OE::H2A.Z. For (C) and (D), letters indicate statistically significant curve groups after Kruskal-Wallis testing and Tukey-Kramer correction for multiple testing (p < 0.05).
I349ΔH2A.Z::H2A.Z that shows no sign of sensitivity to H₂O₂ 5 mM or 15 mM (S5A Fig, S5B Fig and S5D Fig).

The ability to produce mycotoxins was finally investigated in all our mutants and compared to their respective wild-type strains. INRA349 wild type can produce high amounts of DON (and its derivative 15-ADON in vitro, [62,63]), a virulence factor that helps the fungus to spread within wheat spikes [67]. We monitored the kinetics of DON+15-ADON production of wild type and mutants for all strains and expressed yields as micrograms of mycotoxins produced per milligram of dry weight of mycelium (Fig 3A). Toxin production could be detected after three days of culture post-inoculation (dpi) in our conditions for both INRA349 wild type and over-expressed mutant I349-OE::H2A.Z. Both strains then produced toxins
sequencing confirmed that Δwild-type, not be rescued. Thus, in order to shed some light on the causes for our observations, INRA349 successfully re-integrated the phenotypes presented above were puzzling. Indeed, although I349 ΔH2A.Z and I349ΔH2A.Z::H2A.Z only after four days, and production remained very low throughout. Nonetheless, DON and 15-ADON production seemed to slightly increase between 12 and 14 days of culture and thereafter in I349ΔH2A.Z::H2A.Z but not in I349ΔH2A.Z, significantly.

Putative intergenic suppressors rescue deleterious phenotypes in I349ΔH2A.Z and I349ΔH2A.Z::H2A.Z strains

The phenotypes presented above were puzzling. Indeed, although I349ΔH2A.Z::H2A.Z had successfully re-integrated H2A.Z at its locus, most deficiencies caused by H2A.Z deletion could not be rescued. Thus, in order to shed some light on the causes for our observations, INRA349 wild-type, ΔH2A.Z, ΔH2A.Z::H2A.Z, and OE::H2A.Z strains were sequenced by Whole Genome Sequencing (WGS) and all subsequent genome sequences scrutinized. As expected, sequencing confirmed that H2A.Z coding sequence was absent from all ΔH2A.Z strains and successfully added back in I349ΔH2A.Z::H2A.Z (S6A Fig). More surprisingly, we discovered in I349ΔH2A.Z one mutation elsewhere in the genome (S6B Fig) and a second one in I349ΔH2A.Z::H2A.Z (S6C Fig; verified by Sanger sequencing, S7 Fig), which we hypothesized to be compensatory mutations considered to have beneficial effects on fitness when a deleterious mutation is present [68].

The mutation discovered in I349ΔH2A.Z is a deletion of a single nucleotide A in position 2015 of its CDS (S5B Fig), causing a frameshift mutation (Ile673fs) in the ORF of FGRAMPH1_01G23597, annotated in the FungiDB database as an essential subunit of the histone deacetylase Rpd3S complex. FGRAMPH1_01G23597 encodes a 1225 amino acid-long protein predicted to contain two PHD-type domains (one of the TNG2 superfamily and one PHD2_PHF12_Rco1 particularly found in the yeast Rpd3s component Rco1p) including their zinc and histone H3 binding sites (Fig 4A) using the NCBI CD-Search [69,70]. Provided corresponding transcripts with such premature termination codon is not degraded by the nonsense-mediated mRNA decay (NMD) pathway, resulting in no protein at all, the frameshift mutation that occurred in I349ΔH2A.Z would have the drastic consequence of removing both PHD domains and creating a much shorter protein of 783 amino acids (Fig 4A and S8 Fig).

Such truncated protein may either be degraded by the unfolded protein response (UPR) pathway or see its functions very likely dramatically impaired. We hypothesize that this mutation could be responsible for rescuing the deleterious effects of H2A.Z deletion, otherwise lethal. Rpd3S is typically involved in the deacetylation of H3K36. Functions of the acetylated vs. methylated H3K36 are multiple and the subject of intense investigation (reviewed in [71]); methylation of H3K36 is typically associated with active transcription. A proposed mechanism is that SET2-mediated methylation of H3K36 towards the 3' end of genes mediates general histone deacetylation through the recruitment of Rpd3s to slow down transcription elongation at the end of genes [72]. In mammalian embryos, it was suggested that deposition of acetylated H2A.Z (a mark of active transcription) and methylated H3K36 are precisely orchestrated to allow fine-tuned expression of developmental genes and prevent spurious transcription [73].
Here, a frameshift mutation in FGRAMPH1_01G23597 with the likely consequence of losing its activity seems to counterbalance to some extent (the strain can survive) the deletion of H2A.Z. A possible explanation for this rescue could be that transcription elongation speed increases at the end of genes to provide partial compensation for the loss of H2A.Z, in other words the loss of the activation counterpart of a fine-tuned process. In this case, however, fine regulation of transcription can no longer be assured and survival is only permitted at high cost.

In the add-back mutant I349ΔH2A.Z::H2A.Z, in addition to the frameshift mutation in FGRAMPH1_01G23597 inherited from its parental strain I349ΔH2A.Z, another mutation was found in the gene FGRAMPH1_01G18675 encoding the 1,691 amino acid-long helicase Swr1p (S6C Fig) i.e., the catalytic subunit of the SWR1 complex involved in the deposition of H2A.Z (see [39] for a review). This mutation caused the exchange of a histidine at position 852 by a proline in I349ΔH2A.Z::H2A.Z. Domain accession numbers in NCBI CDD/SPARCLE database: TNG2 superfamily = CL34876, PHD = CL22851, PHD2_PHF12_Rco1 = CD15534, HAS = PFAM07529, DEXQc_SRCAP = CD18003, SF2_C_SNF = CD18793. Architectures are displayed with SnapGene Viewer 5.0.6.

Fig 4. Domain architecture of proteins affected by compensatory mutations in I349ΔH2A.Z and I349ΔH2A.Z::H2A.Z. (A) Essential subunit of Rpd3S complex (FGRAMPH1_01G23597; 1,225 aa). The annotation “Ile673fs” indicates the position of a frameshift starting at position 673 of the protein in I349ΔH2A.Z (an isoleucine in wild-type). (B) Swr1 (FGRAMPH1_01G18675; 1,691 aa). The annotation “His852Pro” indicates the replacement of a histidine at position 852 by a proline in I349ΔH2A.Z::H2A.Z. Domain accession numbers in NCBI CDD/SPARCLE database: TNG2 superfamily = CL34876, PHD = CL22851, PHD2_PHF12_Rco1 = CD15534, HAS = PFAM07529, DEXQc_SRCAP = CD18003, SF2_C_SNF = CD18793. Architectures are displayed with SnapGene Viewer 5.0.6.

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Compensatory mutations with varying effects are obtained when \( H2A.Z \) is deleted in different genetic background

Compensatory mutations are defined as beneficial in a particularly deleterious context, and are otherwise undesirable as they usually come as the cost to at least some fitness. Since genetic background influences the outcome of compensatory mutations [76], we engineered \( \Delta H2A.Z \) strains in two additional genetic backgrounds of \( F. graminearum \), INRA156 [77] and the reference strain PH-1 [78–80], and successfully obtained four deletion mutants and one mutant in each of the strain, respectively. All \( \Delta H2A.Z \) deletion strains in INRA156 and PH-1 were severely impacted in their development, but with varying degrees of severity (Fig 5A). The mutant strain INRA156\( \Delta H2A.Z \)\#3 shows the most extreme growth defect, all strains considered. The fact that four mutant strains originating from the same parental strain INRA156 exhibit different phenotypes suggests that the suppressor mutations rather than the influence of the

![Fig 5. Growth defects in INRA156 and PH-1 mutants.](https://doi.org/10.1371/journal.pgen.1009125.g005)
genetic background are responsible for the observed phenotypic variations. Regarding response to oxidative and osmotic/ionic stresses, both INRA156 and PH-1 are the most sensitive to H$_2$O$_2$ 15 mM, and to a much lesser extent NaCl 1M, in terms of radial growth (Fig 5B, S9A Fig and S9F Fig). Nonetheless, both NaCl and KCl caused striking discoloration of the mycelium of wild-type strains (Fig 5B). This discoloration can also be observed in ΔH2A.Z strains submitted to stress (Fig 5B). Nonetheless, marked differences in radial growth sensitivity could be observed. While sensitivity hierarchy is identical in 1156ΔH2A.Z#4 and PH-1ΔH2A.Z to the one in their wild-type counterparts (i.e., mostly to H$_2$O$_2$ 15 mM followed by NaCl 1M, S9E Fig and S9G Fig), the growth curves of 1156ΔH2A.Z#1 and 1156ΔH2A.Z#2 are not significantly affected by any of the applied stresses (S9B Fig and S9C Fig). Finally, the effects observed for 1156ΔH2A.Z#3 (S9D Fig) resemble those observed in I349ΔH2A.Z (S2D Fig), with an increased sensitivity to NaCl and KCl but higher resistance to H$_2$O$_2$ stress.

Differences between mutant strain behaviors were also observed regarding the ability to produce asexual spores. Similar to I349ΔH2A.Z, mutant strains in INRA156 and PH-1 backgrounds were affected in their ability to sporulate (Fig 6A and 6B). The intensity of the defect was however moderate for two of them; 1156ΔH2A.Z#2 (Fig 6A, blue curve) and PH-1ΔH2A.Z (Fig 6B, red curve). Since INRA156 and PH-1 show great capability to produce perithecia in vitro, we also investigated this property in their deletion mutants. The heterogeneity of the consequences of deleting H2A.Z is striking (Fig 6C): two mutants (1156ΔH2A.Z#1 and 1156ΔH2A.Z#3) had their production of perithecia completely abolished; two mutants produced micro-perithecia (1156ΔH2A.Z#4 and PH-1ΔH2A.Z) or not viable ascosores (respectively); one mutant (1156ΔH2A.Z#2) had retained full capability to produce full-size perithecia (although somehow in lesser numbers) that could produce viable spores. Phenotype heterogeneity is also illustrated with the production of the mycotoxins DON and 15-ADON (Fig 3B). Nearly abolished in 1156ΔH2A.Z#3, 1156ΔH2A.Z#4, and PH-1ΔH2A.Z (similar to the situation in I349ΔH2A.Z), toxin production is heavily stimulated in 1156ΔH2A.Z#1, and not significantly affected in 1156ΔH2A.Z#2 (Fig 3C).

We investigated the occurrence of potential compensatory mutations in all ΔH2A.Z strains in both INRA156 and PH-1 background, similarly to the analysis we performed for INRA349 and its derived mutants. Putative compensatory mutations were detected in the two genes previously identified, namely FGRAMPH1_01G23597 and FGRAMPH1_01G18675, plus eight other genes in the five ΔH2A.Z mutants obtained in INRA156 and PH-1 backgrounds (Table 1). Strikingly, most genes are annotated as to encode proteins directly involved in chromatin remodeling in the reference functional database FungiDB [81,82].

In three instances, the same gene is affected in two ΔH2A.Z mutants from two different parental strains, albeit not with the exact same mutation. The first case regards I349ΔH2A.Z and 1156ΔH2A.Z#4 that each carry a version of a frameshift mutation in the essential subunit of the histone deacetylase Rpd3S complex encoded by FGRAMPH1_01G23597. As evoked earlier, frameshifts creating non-sense mutations may result in the absence of the protein rather than the presence of a truncated one due to the activation of the NMD or the UPR pathways. In 1156ΔH2A.Z#4, valine 1,159 becomes the site of a frameshift mutation. This mutation occurs towards the end of the protein and modifies profoundly its primary sequence with the likely consequence of losing the ability to undergo the conformational changes necessary to its proper activity [83]; it does not affect, at least directly, the functional domains of the protein as it can be the case in I349ΔH2A.Z (Fig 4A). Anecdotally, the latter is more severely affected in its growth and response to stress than 1156ΔH2A.Z#4 (see S10 Fig for a visual summary), which suggest a more targeted rescue of H2A.Z deletion that does not (or not so much) compromises other H2A.Z-independent functions. Such differences in phenotypes between I349ΔH2A.Z and 1156ΔH2A.Z#4 are remarkable since they both carry a severe mutation on
Fig 6. Asexual sporulation and formation of perithecia in INRA156 and PH-1 ΔH2A.Z mutants. (A). Fitted kinetics of sporulation for INRA156 wild-type and four ΔH2A.Z mutants. Black = wild-type; red = I156ΔH2A.Z#1; blue = I156ΔH2A.Z#2; purple = I156ΔH2A.Z#3; green = I156ΔH2A.Z#4. (B). Fitted kinetics of sporulation for PH-1 wild-type and PH-1ΔH2A.Z mutants. Black = wild-type; red = PH-1ΔH2A.Z. Letters indicate statistically significant curve groups after Kruskal-Wallis testing and Tukey-Kramer correction for multiple testing (p < 0.05). (C). Formation of perithecia on carrot agar by INRA156 and PH-1 wild-type and corresponding ΔH2A.Z mutants. Top lane: macroscopic view; bottom lane: microscopic view (x40). Pictures were taken four days after induction of sexual differentiation.

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the same gene, albeit not exactly the same one. While phenotypic differences may be attributed to the different genetic backgrounds, an alternative hypothesis is that both events affecting FGRAMPH1_01G23597 did not lead to null loss-of-function in both cases (or any), the alternatives being a leaky loss-of-function (with some functionality preserved) or a gain-of-function.

The second case of genes independently mutated regards I349ΔH2A.Z.ΔH2A.Z.

Z::H2A.

Z and I156ΔH2A.

Z#2 that each harbor mutated SWR1 (FGRAMPH1_01G18675), in addition to other mutations (Table 1). In I156ΔH2A.Z#2, leucine 850 is entirely removed, a mutated site located only two amino acids upstream the His852Pro found in I349ΔH2A.Z::H2A.

Z and within the ATP-binding site of the DEAD box helicase-containing domain (Fig 4B). The activity of mutated FgSwr1, involved in the incorporation of H2A.Z in nucleosomes, is here very likely affected. Another mutation was found in the same mutant I156ΔH2A.Z#2: proline 17 (position 16 in the classical histone numbering since the initial methionine is cleaved off) histone H3 (FGRAMPH1_01G14931) is here replaced by a threonine. Proline isomerization influences protein secondary structure, and protein isomerases are found in many chromatin remodeling complexes [84]. Their exact role is however unclear. In histone H3, interplay between histone lysine methylation and proline isomerization has been demonstrated in S. cerevisiae [85]; for example, proline 38 was found to be necessary for H3K36 methylation. However, the potential function of H3P16 isomerization remains to be uncovered. As mentioned above, Swr1 is a major actor of the deposition of the dimer H2A.Z-H2B to replace a dimer H2A-H2B, notably via interactions with (H3-H4).2. Mutations in both Swr1 and H3 may prevent protein interaction and subsequent removal of H2A-H2B, thus stabilizing durably the histone octamer in the absence of H2A.Z and rescuing part of the deficiency. Among all ΔH2A.Z

| Parental strain | Mutant | Gene ID | Description | Coding-region change | Amino Acid change | Ortholog essential in N. crassa? |
|-----------------|--------|---------|-------------|----------------------|------------------|-----------------------------|
| INRA349         | ΔH2A.Z | FGRAMPH1_01G23597 | essential subunit of the histone deacetylase rpd3s complex | 2015delA | Ile673fs | No |
| INRA349 ΔH2A.Z  | ΔH2A.Z::H2A.Z | FGRAMPH1_01G23597 | essential subunit of the histone deacetylase rpd3s complex | 2015delA | Ile673fs | No |
|                 |        | FGRAMPH1_01G18675 | helicase swr1 | 2555A>C | His852Pro | No |
| INRA156         | ΔH2A.Z#1 | FGRAMPH1_01G16577 | oxidoreductase yusz | 262G>T | Asp88Tyr | No |
|                 |        | FGRAMPH1_01G18925 | histone demethylase jarid1 | 239_240insA | Val81fs | No |
| ΔH2A.Z#2        | FGRAMPH1_01G14931 | Histone H3 | 49C>A | Pro17Thr | Probably essential |
|                 |        | FGRAMPH1_01G18675 | helicase swr1 | 2545_2547delCTT | Leu850del | No |
| ΔH2A.Z#3        | FGRAMPH1_01G03975 | hypothetical protein | 82A>G;100G>A | Thr28Ala; Glu34Lys | n/a |
|                 |        | FGRAMPH1_01G11173 | histone-lysine n-methyltransferase ashl1 | 1609C>T | His537Tyr | Probably essential |
|                 |        | FGRAMPH1_01G26173 | Transcription factor | 741_742insA | Ala248fs | No |
| PH-1            | ΔH2A.Z | FGRAMPH1_01G23597 | essential subunit of the histone deacetylase rpd3s complex | 3474_3475delTG | Val1159fs | No |
|                 |        | FGRAMPH1_01G27197 | Hypothetical protein | 1432C>T | Arg478* | No |

* Source: FungiDB Release 46
**The Neurospora Genome Project [115]; n/a indicates no ortholog identified; identical genes with mutations in more than one genetic background are colored with the same shade of grey
mutants in our set, I156ΔH2A.Z#2 certainly appears as the one in which the deleterious effects of deleting H2A.Z are best rescued.

The last case of genes repeatedly but independently mutated concerns I156ΔH2A.Z#3 and PH-1ΔH2A.Z that each harbor mutated FGRAMPH1_01G03975 (Table 1), encoding a small hypothetical protein with no known conserved domain. Considering that this gene is found immediately downstream and in the vicinity of FgH2A.Z, the mutations observed may be consequences of such proximity during knock-out assays. Nonetheless, other loci are also affected by secondary mutations in both I156ΔH2A.Z#3 and PH-1ΔH2A.Z. In the latter, the CDS of FGRAMPH1_01G27197 is interrupted at arginine 478 (out of 744 amino acids total) by a stop codon removing more than 35% of the protein (Table 1). Proline and alanine-rich (14% and 12%, respectively), its product is annotated as “hypothetical protein” (FungiDB v46) and no clear similarity to known features could shed any light on its possible function. In I156ΔH2A.Z#3, no less than three genes in addition to FGRAMPH1_01G03975 were affected by secondary mutations: FGRAMPH1_01G26683 encoding a hypothetical protein (Pro136Ser), FGRAMPH1_01G26173 encoding a transcription factor (Ala248fs), and FGRAMPH1_01G11173 encoding the histone-lysine methyltransferase Ash1l (Table 1 and S11 Fig). Although its function is unknown, a domain search in FGRAMPH1_01G26683 found a conserved Clr5 domain at the N-terminal end (S11A Fig); in fission yeast, Clr5 is involved in chromatin-mediated silencing and is in the same functional pathway as the histone deacetylase Clr6 (Rpd3). FGRAMPH1_01G26173 is a transcription factor containing a bZIP_P_ATF2 domain (S11B Fig) with a leucine zipper-type dimer interface followed by a proline-alanine-rich region of unknown function. The frameshift mutation occurs in the middle of this region. Finally, the histone methyltransferase Ash1l (SET2, FGRAMPH1_01G11173) contains a SET2_ASH1L domain (S11C Fig) that can bind H3K36 to add methyl groups using co-factor S-adenosyl methionine [86]. The mutation His537Tyr occurs within the active site, likely impeding the binding of S-adenosyl methionine [87]. Remarkably, among all obtained mutants, I156ΔH2A.Z#3 is the one most affected with secondary mutations (hitting four genes) and also the one with the most drastic phenotypes.

The last mutant, I156ΔH2A.Z#1, carries secondary mutations in two genes not affected in any of the other mutants: FGRAMPH1_01G16577 encoding the oxidoreductase Yusz, and FGRAMPH1_01G18925 encoding the histone demethylase Jarid1 (Table 1, and S11D Fig and S9E Fig). FgYusz is a short-chain dehydrogenase/reductase (SDR) with a structurally conserved Rossmann fold (alpha/beta folding pattern with a central beta-sheet; [88–91]). Its function in *F. graminearum* is currently unknown. The mutation Asp88Tyr is located within the NADP-binding site, thus with potential consequences in the activity of the protein (S11D Fig). FgYusz is found highly expressed in conidia, perithecia, and during wheat infection but not in vegetative hyphae [56,58,92]. Our observations of asexual sporulation and perithecia formation being affected whereas radial growth is less severely impaired supports a role in conidiation and sexual reproduction. Regarding the histone demethylase Jarid1 (Kdm5), involved in demethylating H3K4 [93], a single nucleotide insertion has dramatic consequences. A frameshift at the N-terminal end of the protein truncates the protein down to 114 aa with a completely modified sequence from position 81 to 114 (S11E Fig), completely removing all functional domains and binding sites necessary for proper function [94]. Incidentally, I156ΔH2A.Z#1 seems to cope better with stress during vegetative growth, including H2O2 15 mM, than the wild type parent and produces much higher levels of toxins (Figs 3 and 5). This observation is consistent with DON and 15-ADON production being part of *F. graminearum* stress response [66]. FgKdm5 has been recently characterized in *F. graminearum* and was shown to be a positive regulator of secondary metabolism, in a demethylation-independent manner for four out of five secondary metabolites tested including DON [50]. Similarly, in
Aspergillus nidulans, a histone demethylation-independent role of KdmB (FgKdm5) in activating some but not all genes involved in the production of secondary metabolites has been proposed [95]. In the present study, losing the canonical FgKdm5 did not prevent the activation of toxin production, suggesting the detected mutation did not cause of null loss-of-function, or the intervention of another Kdm5-independent pathway stimulated toxin production. In F. fujikuroi the homolog FfKdm5 balances H3K4me3 levels and affects secondary metabolism, hyphal growth and sporulation [96].

As a whole, our observations may suggest that deleting H2A.Z from the genome of F. graminearum causes particularly deleterious effects that would challenge viability in the absence of compensatory mutations.

H2A.Z appears essential also in another Fusarium spp.

To explore the possible essentiality of H2A.Z in other Fusarium spp., attempts to delete H2A.Z were performed in the distantly related rice pathogen Fusarium fujikuroi [97]. F. fujikuroi is a rice pathogen and well known for the production of gibberellins, phytohormones that are produced during the infection and the causative agent for the typical symptoms of the bakanae disease, i.e. chlorotic, slender and etiolated rice stalks [98]. H2A.Z was identified in the F. fujikuroi wild-type strains IMI58289 (FFUJ_01849), B14 (FFB14_11687) and E282 (FFE2_11549) by determining FgH2A.Z orthologs using QuartetS [99], and here referred to as FfH2A.Z. To analyze the function of FfH2A.Z, a gene replacement cassette (pΔffH2A.Z) was generated and used for targeted deletion of FfH2A.Z in all three wild-type backgrounds (S12A Fig). In total, 15 hygromycin-resistant transformants were gained for IMI58289, all heterokaryons of nuclei showing homologous integration events mixed with wild-type ones. However, attempts to generate hygromycin-resistant homokaryotic mutants failed as not one single mutant obtained during the process showed absence of FfH2A.Z (S12B Fig). Similarly, attempts to delete H2A.Z in B14 and E282 resulted in several hygromycin-resistant transformants showing homologous integration events. For three independent transformants each, attempts to generate homokaryotic mutants by single spore isolation were performed. Again, no homokaryotic mutant showing absence of FfH2A.Z was obtained during three repeated rounds of single spore isolation (S12B Fig). Next, we tried to generate homokaryotic mutants by performing protoplast isolation in all three different wild-type backgrounds. Again, not a single hygromycin-resistant homokaryotic strain deleted for FfH2A.Z was obtained (S12B Fig), suggesting that removal of H2A.Z is also lethal in F. fujikuroi.

Conclusion

Histone variant H2A.Z is found ubiquitously in fungi, plant and animal species. In the present study, we conducted attempts to knock out H2A.Z from the F. fujikuroi and F. graminearum genome in three different strains each. While not one single homokaryotic deletion mutant was recovered in all three F. fujikuroi backgrounds, a total of six homokaryotic H2A.Z deletion mutants were obtained in the case of F. graminearum. Notably, all six exhibited heterogenous phenotypes. Moreover, adding-back H2A.Z in I349AΔH2A.Z did not restore wild-type phenotype. We found that each one of F. graminearum genetically modified strain we engineered contained one or more secondary mutations elsewhere in the genome, not present in the wild-type parental strain. Thus, we propose that H2A.Z is essential in both F. fujikuroi and F. graminearum. In the latter, the occurrence of compensatory mutations rescues lethality.

In our study, the six H2A.Z deletion mutants obtained in three different F. graminearum genetic backgrounds harbor different mutations, some of them in the same genes. The fact
that independent repetitions of knock-out experiments often lead to secondary mutations in the same gene has been previously observed in yeast, a phenomenon referred to as parallel evolution [100]. The numerous failed attempts we faced to obtain ΔH2A.Z mutants suggest that there are only a few suitable mutations that could be selected. On the evolutionary scale, the number of combinations that are sufficiently fit to persist may be limited, in line with the proposition from Davis and Colleagues (2007). Such mutations, indeed, usually occur for proteins that are in the same functional module [101]. In *F. graminearum*, an example of the suppression of growth defects provoked by the deletion of the kinase FgPrp4, involved in the spliceosome, has been evidenced and involves spontaneous mutation in *FgSAD1* involved in premRNA complex assembly [102]. In our study, all mutants were affected in at least one known protein involved in chromatin remodeling, with the exception of PH-1ΔH2A.Z for which no function could be proposed for any of the protein containing secondary mutations. Altogether, the critical relationship between H3K36me and H2A.Z is largely highlighted here. The flexible changes of the regulatory networks around these marks reveal the power of network re-wiring to maintain a critical level of transcriptional balance. Some of the genetic innovations that arose from our experiments seem to give better results than others in the light of the traits tested. The overall fitness of our mutant strains to survive and thrive under actual conditions remains to be explored.

The mark H3K4me2/3 has also been underlined as in the same functional network as H2A.Z in our mutant II56ΔH2A.Z#1 defective in the histone demethylase Jarid1 (FgKdm5), which produces much higher levels of toxin than the wild-type. This observation is consistent with findings in *F. fujikuroi* in which methylated H3K4 is deposited by SET1 and positively regulates, among other traits, secondary metabolism [96]. In this mutant, active transcription may be promoted by maintaining methylated H3K4.

As a whole, in the present study, the observed phenotypes are the result of interactions between two or more mutations, under the influence of the genetic background. One feature emerges, lack of H2A.Z is likely lethal in both fusaria investigated here, *i.e.*, *F. fujikuroi* and *F. graminearum*. In the latter, however, its extraordinary plasticity allows compensation within the short time frame of a lab transformation. Our study highlights peculiar links between H2A.Z and H3K36/H3K4, advancing the search for H2A.Z functional network.

**Materials and methods**

**Fungal strains and culture media**

Wild-type *F. graminearum* strains INRA156 (INRA-MycSA collection), INRA349 (CBS185.32; CBS-KNAW Collection, the Netherlands), INRA812 (PH-1 strain FGSC9073; Fungal Genetics Stock Center, the USA) were propagated on Potato Dextrose Agar (PDA; DIFCO, France) plates at 25˚C. INRA156 can efficiently undergo sexual reproduction but produces only moderate to low levels of DON and 15-ADON, whereas INRA349 always fails to produce perithecia but consistently produces very high levels of DON and 15-ADON; INRA812 is the sequenced reference strain widely used in *Fusarium* related studies (can reproduce sexually and produces only very low levels of DON and 15-ADON). Radial growth assays were performed on complete medium [103] supplemented prior solidification with NaCl, KCl, H₂O₂, or not supplemented. When needed, conidia were prepared by inoculating agar plugs in CMC medium (15 g/L carboxymethyl cellulose, 1 g/L yeast extract, 0.5 g/L MgSO₄·7H₂O, 1 g/L NH₄NO₃, 1 g/L KH₂PO₄) [104], incubating at 150 rpm and 25˚C for three to five days, and harvesting by filtration through Sefar Nitex 03–100 (100 μm, SEFAR AG—Switzerland). Toxin production was measured in liquid synthetic medium [105]. For spore germination assays, glucose in MS medium was replaced by sucrose.
Wild-type strains of *F. fujikuroi* IMI58289 (Commonwealth Mycological Institute, Kew, UK), *F. fujikuroi* B14 (S.-H. Yun, Korea), *F. fujikuroi* E282 (S. Tonti, University Bologna, Italy) were used as parental strains for deletion experiments. For protoplasting, relevant strains were grown for three days in 100 mL Darken medium [106] at 30˚C and 180 rpm in the dark. A 500 µL aliquot was then used for inoculation of 100 mL liquid synthetic ICI medium [107] containing 10 g/L fructose and 1g/L NH₄SO₄ as sole carbon and nitrogen sources, respectively. For verification of ΔH2A.Z mutants and DNA isolation, strains were grown for three days on solid complete medium (CM) [108] covered with cellophane sheets (Folia Bringmann) at 30˚C in darkness. Where applicable, plates were supplemented with 100 ppm hygromycin B. For conidia production, *F. fujikuroi* strains were grown for seven days on solid V8 medium (20% v/v vegetable juice, Campbell Food, Puurs, Belgium) at 20˚C and 16/8 light-dark cycles. Cultivation of *S. cerevisiae* was performed as previously described [109].

**Identification of FgH2A.Z**

We used a Hidden Markov Model-based approach [110] to identify proteins carrying a domain characteristic of the C-terminal end of histone H2A (PFAM accession number PF16211) in the reference proteome of *F. graminearum* PH-1 [78–80] available at FungiDB [81,82]. Putative identification was then assigned by sequence similarity with the reference protein database (ref_seq_protein_v5) at NCBI using BLASTP [55,111]. Genes structures and sequences, protein sequences, and transcript levels were retrieved from FungiDB [81,82]. Protein alignment was performed and displayed with CLC Workbench 10.0.

**Generation of gene deletion, add-back, and over-expression mutants**

Wild-type *F. graminearum* strains INRA156, INRA349, and PH-1 were used as parental strains to engineer H2A.Z deletion mutants. Primer sequences and amplification conditions used are provided in S1 Table. Deletion strategy using the split-marker approach [112] is represented S2 Fig. Upstream and downstream flanking regions of FgH2A.Z (gene ID FGRAMP H1_01G03973) were amplified by PCR (primer pairs 3r-3UTR-H2AZ-R / 3UTR-H2AZ-HY-F, and 5UTR-H2AZ-GRO-R / 5f-UTR-H2AZ-F, respectively) from wild-type INRA349 genomic DNA. The hygromycin-resistance cassette hygroR, containing *hph* gene under the control of *N. crassa* pCPC1 and *A. nidulans* *tTrpC*, was amplified by PCR using primer pair neoHY-8-finF / neoHY-1-debutR and pBSK(-) NeoHygroR plasmid as template according to a previously described procedure [113]. Fragment assembly was achieved by multiple recombination of overlapping sequences in *S. cerevisiae* strain FY1679 (genotype MATa/MATα, ura3-52/ura3-52, trp1Δ63/TRP1, leu2Δ/LEU2, his3A200/HIS3, GAL2/GAL2) transformed with the three PCR products and the pRS426 plasmid digested with BamHI and HindIII [114]. SC-Ura medium containing 0.67% yeast nitrogen base without amino acids was used for selection. A second round of PCR was carried out using successfully transformed FY1679 DNA as template to amplify two overlapping fragments (primer pairs 3UTR-H2AZ-N-R / NP_SplitHY_R, and NP_SplitGRO_F / 5UTR-H2AZ-N-F) to be used for transformation into *F. graminearum* for the targeted replacement of H2A.Z with the hygromycin resistance cassette (hygroR). PCR products were purified and transformed into protoplasts of INRA156, INRA349, and PH-1, prepared according to a previously published protocol [113]. For add-back experiment, H2A.Z CDS fused to *tTrpC* terminator and followed by a geneticin-resistance cassette, genetR as previously described [113], (assembled in yeast similarly to the deletion experiments) was used for targeted replacement of the integrated hygroR cassette in INRA349AH2A.Z (S3A Fig and S1 Table). For over-expression experiments, wild-type H2A.Z CDS in INRA349 was replaced by FgH2A.Z CDS fused to *A. nidulans* promoter *pGPD* and preceded by the above described
HygroR cassette (S3B Fig and S1 Table). All selected transformants were purified by two rounds of monoconidial isolation.

The plasmid for H2A.Z deletion in *F. fujikuroi* (IMI58289, B14 and E282) was generated using yeast recombinational cloning [109,115]. All primers used for polymerase chain reaction (PCR) were obtained from Sigma-Aldrich GmbH. Deletion strategy and respective primers for H2A.Z deletion in the different wild-type strains are displayed in S12 Fig as well as listed in S2 Table. Briefly, the upstream (5’) and downstream (3’) sequences of *FfH2A.Z* were amplified from *F. fujikuroi* IMI58289 wild-type genomic DNA using the following primer pairs: FfH2A.Z_5F // FfH2A.Z_5R for upstream, FfH2A.Z_3F // FfH2A.Z_3R for downstream regions. For the deletion construct, hygromycin B was used as a selection marker. The hph resistance cassette was amplified from pCSN44 [116] with the primer pair Hph_F // Hph_R. Transformation of *F. fujikuroi* (IMI58289, B14 and E282) was performed as previously described [97,117]. For this, the deletion fragments were amplified from pΔfH2A.Z with the primer pairs FfH2A.Z_5F // Ff_H2A.Z_3R, using a proof-reading polymerase (Q5-polymerase, New England Biolabs). Transformed protoplasts were regenerated as described [118]. Regeneration media contained 100 ppm hygromycin B. Homologous recombination events were verified with the primer pairs dia_FfH2A.Z_F // pCSN44_trpC_T for the upstream part and dia_FfH2A.Z_R // pCSN44_trpC_P2 for the downstream part (S12 Fig). Presence of the native wild-type gene was checked with the primers FfH2A.Z_WTDiaF // FfH2A.Z_WTDiaR.

**Radial growth assays**

A total of 100 spores or a plug (3 mm in diameter, taken from the periphery of a seven-day-old CM culture plate) were cultured at 25˚C in darkness on 94 x 16 mm plates containing CM, or CM with 1M NaCl / KCl, or CM with 5 mM / 15 mM H2O2. Experiments were done in triplicate. Growth area of individual Petri dishes was analyzed by ImageJ 1.8.0 image processing program [119].

**Conidiation and germination rate assays**

For conidiation assays, one mycelial plug (8 mm in diameter) of each strain, taken from the periphery of a 7-day old colony on PDA plates was inserted in a 50 mL-filter screw cap conical tube containing 10 mL of CMC medium, and incubated for up to eight days at 25˚C and 180 rpm in darkness in an Infors Multitron (INFORS-HT). Spores were counted in a Thoma cell counting chamber under microscope every 24 hours (three independent counts per sample). For germination assays, 10^6 five-day old spores were incubated in 10 mL MS sucrose liquid medium at 25˚C in darkness for up to eight hours. Germination rates were counted under microscope every hour, according to a previously published procedure [63]. All experiments were performed with three independent biological replicates.

**Sexual reproduction assays**

Self-fertilization were performed on carrot agar plates as previously described [120]. Perithecia were observed under microscope two days after sexual induction. Ascospores were harvested by placing the carrot agar plates upside down and allowing the mature perithecia to discharge ascospores on petri dish covers. Ascospores were collected by adding 1 mL of sterile MilliQ water and observed under microscope. A volume of 10 μL of each suspension was inoculated in liquid CM to check whether ascospores could grow or not. Experiments were done in triplicate.
Type B trichothecene analysis
Eight milliliters of MS liquid medium [105] were inoculated with 8x10^4 conidia prepared in CMC medium and incubated at 25°C in the dark for up to 20 days. TCTB were extracted and analyzed by HPLC-DAD according to a previously published protocol [113].

Whole genome sequencing and data analysis
Genomic DNA was extracted from 50 mg of freeze-dried mycelium, from all wild-type and mutant strains, as previously described [121]. Libraries were prepared from 500 ng of gDNA using the Westburg NGS DNA Library Prep Kit for Illumina (cat. # WB9024, Westburg, The Netherlands) following manufacturer’s instructions. Sequencing in paired mode, 2x100 bp, was performed by the GenomEast platform, a member of the ‘France Génomique’ consortium (ANR-10-INBS-0009) on an Illumina HiSeq 4000 (reads were deposited at SRA under the Bio-Project accession number PRJNA580372). Raw reads were pre-processed with Trimmomatic v0.39 [122]. The software CLC Genomics Workbench 10 (QIAGEN Bioinformatics, Denmark) was used for all subsequent bioinformatic analyses. Briefly, high-quality read pairs were mapped (including a re-alignment step) onto F. graminearum PH-1 reference genome. Coverage analysis was performed to confirm the correct insertion of the deletion/complementation cassette in all mutants. Variants were called (ploidy = 1 and variant probability = 0.9), and those found in wild-type were subtracted from their corresponding mutants. Each remaining variant was manually curated to ensure no false positive was retained. Fragments including compensatory mutations identified by whole genome sequencing of ΔH2A.Z, ΔH2A.Z::H2AZ and their corresponding wild-type strains were amplified by PCR. Products were sent to GENEWIZ for Sanger sequencing. Fragments were sequenced by two primers from the direction of forward and reverse, respectively (S1 Table). The software CLC Genomics Workbench 10 (QIAGEN Bioinformatics, Denmark) was used for sequence alignment and trace examination.

Statistical analyses
For growth, sporulation, and germination assays, profiles were obtained by curve fitting through all points and differences statistically analyzed by Kruskal-Wallis test combined with Tukey-Kramer correction for multiple testing using Matlab R2019a (Mathworks). The statistical threshold was set at p = 0.05 throughout.

Supporting information
S1 Text. Additional materials and methods, and additional references.
(DOCX)

S1 File. BLASTP report for query translated protein FGRAMPH1_01T26109.
(PDF)

S2 File. BLASTP report for query translated protein FGRAMPH1_01T03973.
(PDF)

S1 Fig. Schematic representations of F. graminearum H2A (FGRAMPH1_01G26109) vs. H2AZ (FGRAMPH1_01G03973) gene and protein sequences. (A) Genomic sequences. Blue boxes are exons (e); green boxes are introns; black lines are UTRs. Translation start (ATG) and stop sites are indicated with orange and yellow arrows, respectively. Scaling key is provided in grey. (B). FgH2A and FgH2A.Z expression levels (in FPKM) previously measured by Zhao and Colleagues [1] in asexual spores (black bars) or actively growing mycelium (white bars). Displayed values are means of three replicates and error bars are standard deviations.
(C). Protein alignment of FgH2A and FgH2A.Z. Residues were colored according to Rasmol scheme. Red boxes indicate domains essential for H2A.Z function (vs. H2A) according to Suto et al. [2]. All datasets can be viewed at FungiDB [3,4] (see ‘Additional References’ in S1 Text).

**S2 Fig.** Split-marker approach used for the deletion of H2A.Z in *F. graminearum*. (A) The upstream and downstream flanking regions of H2A.Z, HygroR cassette were amplified from wild type *F. graminearum* genomic DNA and plasmid DNA of pBlueScriptSK(-)_NeoHygroR, respectively. (B) Fragment assembly in yeast. The three fragments were assembled by plasmid pRS426. (C) Extraction of yeast genomic DNA and amplification of fragments for split-marker method in *F. graminearum*. (D) Transformation in *F. graminearum*. H2A.Z was replaced by HygroR cassette.

**S3 Fig.** I349 deletion mutant screening by PCR and Southern-Blot (see ‘Additional Materials and Methods’ in S1 Text). (A) Schematic representation of the loci amplified. Primers matching the upstream and downstream flanking regions of H2A.Z were used to amplify a 1,588 bp- or 2,717 bp-long fragments from wild-type INRA349 gDNA or ΔH2A.Z mutant gDNA, respectively. (B) Result of the agarose gel electrophoresis of the fragments obtained from S3A Fig. Lane 1: ladder; lane 2: INRA349 WT; lane 3: I349ΔH2A.Z. (C) Schematic representation of the Southern blot strategy used. Digestion with NdeI of gDNA extracted from I349 wild-type or ΔH2A.Z leads to fragments of 7,687 bp and 3,847 bp in size, respectively. A probe marked with digoxygenin and matching the upstream region of H2A.Z CDS as well as part of its 5' end was then synthesized and used to reveal the two digestion fragments obtained on X-ray films (as displayed in (D); Lane 1: INRA349 WT; lane 2: I349ΔH2A.Z. WT = INRA349 WT; Δ = I349ΔH2A.Z.

**S4 Fig.** Schematic representation of the split-marker strategy used to add back H2A.Z to INRA349ΔH2A.Z at its original locus (A), or place it under the control of the strong constitutive promoter pGPD (B). (C) Rq expression levels for H2A.Z in INRA349OE::H2A.Z (grey bar; Rq = 7.6) and INRA349ΔH2A.Z::H2A.Z (striped bar; Rq = 0.978) relative to wild type (black bar; Rq = 1), measured by RT-qPCR. Displayed values are means of three replicates, error bars are standard deviations. The star indicates significant difference with p < 0.01.

**S5 Fig.** Effect of abiotic stress on radial growth. (A) 3 dpi and (B) 6 dpi radial growth in I349ΔH2A.Z (light grey), I349ΔH2A.Z::H2A.Z (stripes), and I349OE::H2A.Z (dark grey) expressed relative to growth of wild-type in not supplemented CM medium (black solid line marking 100% of growth). (C) and (D) Fitted radial growth kinetics followed for six days for I349 OE::H2A.Z (C) and I349ΔH2A.Z::H2A.Z (D). Black = CM; red = CM + NaCl 1M; blue = CM + KCl 1M; purple = CM + H2O2 5 mM; green = CM + H2O2 15 mM. In all panels, letters indicate statistically significant curve groups after Kruskal-Wallis testing and Tukey-Kramer correction for multiple testing (p < 0.05).

**S6 Fig.** Visualization of deep sequencing results at H2A.Z for INRA349 wild-type, its deletion mutant I349ΔH2A.Z, and the add-back I349ΔH2A.Z::H2A.Z. (A) JBrowse screenshot of read coverages per base. (B) and (C) Mutations detected elsewhere in the genomes of I349ΔH2A.Z (B) and I349ΔH2A.Z::H2A.Z (C).
S7 Fig. Snapshot of Sanger sequencing results of *F. graminearum* wild-type and mutant strains. Nucleotides in black are nucleotides that are different.

(TIFF)

S8 Fig. Consequences of the Ile673fs mutation on the primary sequence of the protein encoded by FGRAMPH1_01G23597 in I349ΔH2A.Z. Protein sequence alignment of wild-type (WT) and mutated (ΔH2A.Z) FGRAMPH1_01G23591. Only the C-terminal end of the protein, containing the mutation is depicted. Pink boxes highlight differences between sequences.

(TIFF)

S9 Fig. Fitted radial growth kinetics followed for six days for INRA156 wild-type (A) and its four ΔH2A.Z mutants (B) to (E), and PH-1 wild-type (F) and its ΔH2A.Z mutant (G). Black = CM; red = CM + NaCl 1M; blue = CM + KCl 1M; purple = CM + H2O2 5 mM; green = CM + H2O2 15 mM. Letters indicate statistically significant curve groups after Kruskal-Wallis testing and Tukey-Kramer correction for multiple testing (*p* < 0.05).

(TIFF)

S10 Fig. Visual comparison of developmental defects in the deletion mutants I349ΔH2A.Z and I156ΔH2A.Z#4 that both carry a frameshift mutation in FGRAMPH1_01G23597. (A) Fitted radial growth kinetics followed for six days. (B) Fitted kinetics of sporulation. (A) and (B) Solid black = wild-type I349; dashed black = wild-type I156; solid red = I349ΔH2A.Z; dashed red = I156ΔH2A.Z#4. (C) Effect of NaCl (red bars), KCl (blue bars), H2O2 5 mM (purple bars), and H2O2 15 mM (green bars) after 6 day-long radial growth of INRA349 wild-type (I349WT), INRA156 wild-type (I156WT), and their deletion mutants I349ΔH2A.Z (I349Δ) and I156ΔH2A.Z#4 (I156Δ4). For comparison sake, growth of a given strain is expressed in percentage relative to itself grown in not supplemented CM medium (black bars).

(TIFF)

S11 Fig. Domain architecture of proteins affected by secondary mutations in INRA156 and PH-1 ΔH2A.Z mutants. (A) Protein of unknown function (FGRAMPH1_01G26683; 537 aa) containing a Clr5 domain (PFAM144420). “Pro136Ser” indicates a mutation. (B) Transcription factor (FGRAMPH1_01G26173; 297 aa) containing a bZIP_ATF2 domain (CL21462). “Ala248fs” indicates a mutation. (C) SET2 protein (FGRAMPH1_01G11173; 786 aa) containing a SET_ASH1L domain (CL1917) and an AWS domain (PFAM17907). “His537Tyr” indicates a mutation. (D) Yusz oxidoreductase (FGRAMPH1_01G16577; 292 aa) containing a 17beta-HSD-like_SDR_c domain (CD05374). “Asp88Tyr” indicates a mutation. (E) Jarid1 histone demethylase (FGRAMPH1_01G18925; 1,731 aa) containing various domains (PLU-1 PFAM08429, JmJc PFAM02373, ARID SMART01014, PHD_Ecm5p_Lid2p_like CD15518, JmJN SMART00545, PHD_SF super family CL22851, zf-C5HC2 PFAM02928). “Val81fs” indicates a mutation. Architectures are displayed with SnapGene Viewer 5.0.6.

(TIFF)

S12 Fig. Summary of H2A.Z deletion attempts in *F. fujikuroi*. (A) Schematic representation of the deletion strategy and position of the primers used for PCR validation. (B) Visualization of typical PCR results after three rounds of single-spore isolation. + = ΔffH2A.Z gDNA before single spore isolation; FFWT = *F. fujikuroi* wild-type gDNA. Results are shown for three transformants per strain tested. (C) Visualization of PCR results after protoplast isolation. + = ΔffH2A.Z gDNA before protoplast isolation.

(TIFF)
S1 Table. Primers used in the *F. graminearum* study.
(XLSX)

S2 Table. Primers used in the *F. fujikuroi* study.
(XLSX)

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