GPCR-mediated glucose sensing system regulates light-dependent fungal development and mycotoxin production

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Abstract

Microorganisms sense environmental fluctuations in nutrients and light, coordinating their growth and development accordingly. Despite their critical roles in fungi, only a few G-protein coupled receptors (GPCRs) have been characterized. The Aspergillus nidulans genome encodes 86 putative GPCRs. Here, we characterize a carbon starvation-induced GPCR-mediated glucose sensing mechanism in A. nidulans. This includes two class V (gprH and gprI) and one class VII (gprM) GPCRs, which in response to glucose promote cAMP signalling, germination and hyphal growth, while negatively regulating sexual development in a light-dependent manner. We demonstrate that GprH regulates sexual development via influencing VeA activity, a key light-dependent regulator of fungal morphogenesis and secondary metabolism. We show that GprH and GprM are light-independent negative regulators of sterigmatocystin biosynthesis. Additionally, we reveal the epistatic interactions between the three GPCRs in regulating sexual development and sterigmatocystin production. In conclusion, GprH, GprM and GprI constitute a novel carbon starvation-induced glucose sensing mechanism that functions upstream of cAMP-PKA signalling to regulate fungal development and mycotoxin production.

Author summary

Sensing and responding to the environment is essential for the survival of all fungi. Despite our detailed knowledge of the signalling events that coordinate these adaptations, the extracellular receptors and stimuli that trigger these adaptations remain elusive. G-protein coupled receptors (GPCRs) are the largest class of receptors in fungi and regulate multiple aspects of their biology. Here, we characterized a novel GPCR-mediated glucose...
sensing mechanism in *Aspergillus nidulans*, which influences fungal development and toxin production. This work suggests fungal GPCRs sense glucose and propagate their signals through the cAMP-PKA and light-responsive VeA pathways to promote vegetative growth, while repressing sexual development and toxin production. Then we show how light also impacts on these nutrient sensing mechanisms. Our work reflects the complex and integrated structure of fungal mechanism to perceive different environmental stimuli, and how these mechanisms provide developmental and metabolic versatility to enable fungi to adapt to a fluctuating environment. Fungal GPCRs are poorly conserved in animals and plants, making them relevant antifungal drug targets to intervene in fungal disease and toxin contamination.

**Introduction**

Sensing and responding to fluctuations in the surrounding environment, such as nutrients and light, is crucial to a microorganism’s survival. In eukaryotes, G-protein signalling mechanisms play a vital role in environment sensing, and are typically composed of G-protein coupled receptors (GPCRs), a heterotrimeric G-protein complex consisting of α, β and γ subunits, and a variety of effector proteins [1, 2]. GPCRs communicate changes in the extracellular environment to intracellular G-proteins that initiate and direct signalling events to coordinate the appropriate transcriptional response. In fungi, G-protein signalling regulates cell growth and division, mating, cell-to-cell fusion, morphogenesis, chemotaxis, secondary metabolite production and virulence [3–7]. However, the functions of fungal GPCRs are less well understood and only a few putative receptor-activating ligands have been identified.

*Aspergillus nidulans* has been used in research as a model filamentous ascomycete fungus for more than six decades [8]. It is a saprophytic food spoilage mould that is phylogenetically related to *Aspergillus fumigatus*, the opportunistic pathogen of immunocompromised individuals [9]. The *A. nidulans* genome is predicted to encode 86 putative GPCRs, which are classified according to their structural similarities and putative activating ligands [10, 11]. Sixteen receptors, named GprA-GprP and NopA, constitute nine categories of GPCRs. Finally, 70 *A. nidulans* class X Pth11-like receptors, which promote fungal-plant pathogenic interactions, were identified [10, 12, 13].

Nutritional state and the perception of a sexual partner regulate sexual development. In *A. nidulans*, the formation of sexual structures is promoted by favourable nutrient-rich conditions and inhibited by carbon or amino acid starvation [4, 14]. The GprA and GprB pheromone receptors are required for sexual development, but not vegetative growth [15], while the putative GprD carbon receptor promotes hyphal growth and conidial germination when grown in the presence of glucose [4]. GprD-mediated carbon sensing also acts as a repressor of sexual development, and potentially functions upstream of sex pheromone signalling. But this phenotype is abolished during carbon starvation, suggesting the presence of other repressors of sex under this condition. The putative GprH carbon and amino acid receptor functions upstream of the cAMP-activated Protein Kinase A (PKA) pathway, promoting glucose uptake and hyphal growth, while repressing sexual development, during carbon starvation [16]. Therefore, the GprD and GprH mechanisms appear to operate under distinct environmental conditions and repressing sexual development.

Light also influences fungal developmental decisions [17–19], with *A. nidulans* predominantly producing asexual conidia in the light and sexual fruiting bodies (cleistothecia) in the dark [20–22]. The Velvet family proteins, VeB, VeC and VosA transcription factors, plus the
VeA global regulator gene, are light-dependent regulators, which interact with the LaeA methyltransferase, to regulate fungal development and secondary metabolism [23]. The interaction between VeA and VelB is essential for cleistothecia formation and is established in the cytoplasm. In the dark, the dimer is transported to the nucleus [17], where VeA also interacts with LaeA, affecting sexual development [24]. Additionally, in the dark VelB binds VosA and represses asexual conidiation, while under light, LaeA reduces VosA and VelB levels, triggering asexual sporulation [25].

Development and secondary metabolism are genetically linked in A. nidulans [26, 27]. Aspergillus nidulans produces a variety of secondary metabolites which are toxic to humans and animals, including, sterigmatocystin (ST) [28], the penultimate precursor of aflatoxins produced by related Aspergillus species [29, 30]. The ST biosynthetic gene cluster is regulated by the cluster-specific AflR transcription factor [31], which operates downstream of glucose sensing, G-protein signalling and the cAMP-PKA [32]. Additionally, LaeA also regulates the ST biosynthetic gene cluster in an AflR-dependent manner. The ΔlaeA mutation abolishes aflR expression and subsequently the production of ST and other secondary metabolites [33]. The cross-talk between light and glucose sensing in the coordination of ST production and fungal development is mediated by VeA [34]. Similarly, the ΔveA and ΔvelB mutations that disrupt the formation of Velvet complexes impair ST production [24, 35]. GPCRs play a critical role in perceiving these environmental stimuli and regulating the appropriate signalling pathways [36]. However, the importance of GPCRs and the functional connections between them in regulating secondary metabolism are unknown.

Here, we characterise a novel carbon starvation-induced GPCR mechanism in A. nidulans. This consists of three GPCRs, two class V receptors (gprH and gprI) and a single class VII (gprM) receptor. These GPCRs respond to glucose, promoting cAMP signalling, germination and hyphal growth, while negatively regulating sexual development in a light-dependent manner. We then show that GprH coordinates sexual development via regulating VeA nuclear localisation and activity, a key light-dependent regulator of fungal morphogenesis and secondary metabolism. Additionally, our studies revealed that GprH and M are light-independent negative regulators of sterigmatocystin biosynthesis. Genetic studies showed the epistatic interactions between the three GPCRs in the regulation of sexual development and sterigmatocystin mycotoxin biosynthesis. These studies reflect the complexity and integrated structure of fungal GPCR-mediated mechanisms in coordinating the perception of environmental stimuli with fungal development and secondary metabolism, to achieve optimal adaptations to a changing environment.

Results

The transcriptional regulation of GPCR encoding genes during carbon starvation

The expression of the 16 class I-IX GPCR encoding genes, named gprA-gprP and nopA, was assessed during growth on glucose and carbon starvation (Table 1). This included two class V receptors, gprH and gprI, with similarity to the Dictyostelium discoideum cAMP receptor cAR1 [37]. The previous genome-wide microarray study of the transcriptional response of A. nidulans to carbon starvation showed that class V receptor gprH was transcriptionally induced during carbon starvation, where it regulated glucose uptake, hyphal growth and repressed sexual development [16, 38]. The gene model for the other class V receptor, gprI (AN8348), was found to be inaccurate on chromosome V of the FGSC_A4 genome annotation within FungiDB database [39] and was not predicted to yield a 7-TM containing GPCR. The BLAT alignment of this gene model was also not in agreement with other Aspergillus species and closely
related fungi. Using RNA-sequencing data, a modified *A. nidulans* AN8348_M model was corrected to encode a class V receptor with 7-TM and the Dicty_CAR domains (S1 Fig).

Among the remaining GPCR encoding genes, the *gprA* α-pheromone receptor was induced despite carbon starvation representing a condition repressive of sexual development. The three putative class III carbon receptors *gprC-E* did not show a major increase in expression. Among the three class IV putative nitrogen receptors possessing a PQ-loop repeat *gprF* and *gprJ* were highly repressed, while *gprG* which only is predicted to encode 5-TMs was induced. The class VI *gprK* receptor with the cytoplasmic repressor G-protein signalling domain, and the class IX opsin receptor, were repressed during carbon starvation. Single class VII (*gprM*) and VIII (*gprO*) receptors were induced, however, the latter showed greater variation in expression. Therefore, *gprI* which represented another uncharacterized class V receptor and *gprM* which represented an uncharacterized receptor consistently induced by carbon starvation, were selected to future investigation.

We hypothesized that these three receptors (*gprH, gprI* and *gprM*) may represent a nutrient sensing mechanism that is activated during carbon starvation. To investigate their functions and the potential genetic interactions between these receptors, the Δ*gprH, ΔgprI* and Δ*gprM* null mutant strains were generated in the *A. nidulans* AGB551 wild-type, veA+, background (S1 Table). The expression of the three GPCR encoding genes during carbon starvation in the wild-type strain, and its absence in the mutant strains, was confirmed via RT-qPCR. In the wild-type strain, *gprH* and *gprM* were transcriptionally induced after the shift to carbon starvation, where *gprH* showed the highest level and fastest rate of induction, while *gprI* expression was not detected (Fig 1A), validating the previously observed microarray data.

The dependency of the transcriptional induction of individual GPCRs during carbon starvation on the presence of the other two GPCRs was assessed (Fig 1B). The absence of *gprI* or *gprM* lead to the increased expression of the *gprH* during carbon starvation. Similarly, *gprM*

| Gene name | Gene ID | Class | Domains (No. TMs) | Expression 12 h | Carbon starvation |
|-----------|---------|-------|-------------------|-----------------|------------------|
|           |         |       |                   | Mean  | Std  | Mean  | Std  |   |
| *gprH*    | AN8262  | V     | Secretin-like; Dicty_CAR; Family 2-like (7-TM) | Up    |      | 1.83  | 0.21 | 1.86  | 0.21 |
| *gprI*    | AN8348  | V     | Dicty_CAR; Family 2-like (7-TM) | Absent |      |       |      |   |
| *gprJ*    | AN5720  | IV    | PQ-loop repeat (7-TM) | Down  |       | -1.43 | 0.02 | -1.26 | 0.03 |
| *gprK*    | AN7795  | VI    | GprK-like; RGS (7-TM) | Down  |       | -1.34 | 0.09 | -1.81 | 0.16 |
| *gprM*    | AN6680  | VII   | MG00532-like (7-TM) | Up    |       | 0.49  | 0.07 | 0.72  | 0.05 |
| *gprN*    | AN5508  | VII   | MG00532-like (6-TM) | N/A   |       |       |      |   |
| *gprO*    | AN4932  | VIII  | mPR-like; Haemolysin-III related (7-TM) | Up    |       | 0.81  | 0.21 | 0.97  | 0.38 |
| *gprP*    | AN5151  | VIII  | mPR-like; Haemolysin-III related (7-TM) | N/A   |       |       |      |   |
| *nopA*    | AN3361  | IX    | Bacterial rhodopsin-like (6-TM) | Down  |       | -3.44 | 0.14 | -2.73 | 0.22 |

* GprI gene model corrected to ChrV, antisense, 194574–1957377. Therefore, expression data is absent from array.

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A

\[ \text{gprH} / \text{tubC} \]

\[ \text{gprI} / \text{tubC} \]

\[ \text{gprM} / \text{tubC} \]

WT | ΔgprH
---|---
Glu, St 4h, St 8h | Glu, St 4h, St 8h | Glu, St 4h, St 8h

B

\[ \text{gprH} \]

\[ \text{gprI} \]

\[ \text{gprM} \]

0 | 12

C

[Graph showing cAMP production over time with different conditions: WT, ΔgprH, ΔgprI, ΔgprM, ΔgprHΔgprM, ΔgprMΔgprI, ΔgprHΔgprIΔgprM]
Fig 1. Carbon starvation induced gprH, gprI and gprM expression, which influence cAMP signalling. A-B) Conidia of the wild-type (WT), ΔgprH, ΔgprI and ΔgprM strains were grown in MM with 1% glucose, then subsequently washed and transferred to MM lacking a carbon source for 4 and 8 h (short-term carbon starvation). Steady-state mRNA levels of each individual gpr (gprH, gprI and gprM) were measured by RT-qPCR and normalized with tubC. A) Expression levels of gprH, gprI and gprM in the WT and ΔgprH strains are presented as copy number ratio relative to the reference gene. Statistical analysis was performed using two-way ANOVA (Analysis of Variance) with Bonferroni post-test when compared to the WT strain in each condition (Fig 1). Carbon starvation induced gprM in the WT and strains were grown in MM with 1% glucose, then subsequently washed and transferred to MM lacking a carbon source for 4 and 8 h (short-term carbon starvation). Steady-state mRNA levels of each individual gpr genes during carbon starvation, corresponding to the WT, ΔgprH, ΔgprI and ΔgprM strains. Data presented as fold induction. Grey indicates expression not detected. Glu indicates glucose, St indicates starvation. C) Conidia of the corresponding gpr null mutants and the WT strain were grown in YG for 16 h, then mycelia were washed and incubated in MM lacking a carbon source for 4 h. Glucose at final concentration of 2% was added and incubated for up to 10 min. The results are the average of two biological replicates (with two technical repetitions). Statistical analysis was carried out using two-way ANOVA test with Bonferroni post-test when compared to the WT strain (‘p < 0.05, ** p < 0.01, *** p < 0.001). cAMP levels presented as fmol/μg protein.

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showed increased expression in the ΔgprH mutant after 8 h of carbon starvation. However, no gprI expression was detected in any of the strains. The increased expression of different GPCRs in the absence of others, implies that functional redundancy and compensatory functions may exist among these three GPCRs during carbon starvation.

Identification of GprH, GprI and GprM as a carbon starvation-induced, putative carbon sensing mechanism regulating cAMP signalling

We previously showed that the absence of GprH affected the production of a glucose-induced cAMP burst, glucose uptake and the recovery of growth in starved germlings, suggesting that GprH sensed glucose and functioned upstream of the cAMP-PKA pathway [16]. Accordingly, we quantified intracellular cAMP levels in the ΔgprH, ΔgprI and ΔgprM strains during carbon starvation and after the re-introduction of glucose. In the wild-type strain, the addition of glucose resulted in a burst in cAMP, a response that was absent in both the ΔgprI and ΔgprM strains (Fig 1C). This was reminiscent of the absence of a glucose-induced cAMP burst in the ΔgprH strain. This suggests that GprH, GprI and GprM are all putative carbon sensors functioning upstream of the cAMP-PKA pathway in A. nidulans. Double (ΔgprHΔgprI, ΔgprHΔgprM, ΔgprIΔgprM) and triple (ΔgprHΔgprIΔgprM) mutants were used to investigate the possible genetic interaction between the three receptors in the regulation of cAMP signalling post carbon starvation. The simultaneous absence of multiple GPCRs, in the double or triple null mutants, resulted in a recovery of the burst in cAMP production post the addition of glucose (Fig 1C). This suggests that the absence of multiple GPCRs may activate a compensatory cAMP signalling mechanism.

GprH, GprI and GprM influence conidial germination and hyphal growth in response to glucose

In A. nidulans the cAMP-PKA pathway positively regulates conidial germination in the presence of glucose [40]. Subsequently, we determined if the three putative glucose receptors also influenced conidial germination in the presence of glucose. The ΔgprH strain showed a reduction in germination frequency in comparison to the wild-type strain, suggesting that gprH may positively regulate conidial germination (Fig 2A). No differences in germination frequency were observed for the ΔgprI or ΔgprM strains. The combination of ΔgprH with either ΔgprI or ΔgprM, decreased germination frequency compared to the wild-type strain, to a similar extent as the single ΔgprH strain (Fig 2B). Interestingly, combining the ΔgprI and ΔgprM mutations also decreased conidial germination, a phenotype that was absent in the corresponding single null mutants, suggesting the occurrence of an additive genetic effect and redundancy between the two receptors. However, the combination of all three ΔgprH, ΔgprI and ΔgprM deletions restored germination frequency to wild-type levels. Collectively, this suggests that the GprH signalling mechanism may function in parallel to GprI and GprM, which function in epistasis,
to regulate conidial germination in response to glucose, while the absence of all three receptors may activate a compensatory mechanism.

The cAMP-PKA pathway in *A. nidulans* also positively regulates vegetative hyphal growth [6, 40]. GprH was previously shown to influence glucose uptake and the recovery of growth in
starved germlings [16]. The recovery of hyphal growth after starvation and the re-introduction of glucose was assessed to determine if GprH, GprI and GprM genetically interact during these conditions. As we previously described for the ΔgprH strain, the ΔgprI and ΔgprM strains also displayed a delay in the recovery of growth after the re-introduction of glucose to starved cells (Fig 2C). Analysis of the double and the triple null GPCR mutants unexpectedly revealed that the combination of two, or the three, null GPCR mutations did not impair hyphal growth, and in fact showed increased germling length after the re-introduction of glucose (Fig 2D). This reflects the recovery of glucose-induced cAMP signalling in the simultaneous absence of multiple GPCRs. Taken together, these data suggest that GprH, GprI and GprM represent carbon starvation-induced glucose sensing mechanisms that promote germination and hyphal growth in A. nidulans. However, the absence of multiple receptors activates an unknown compensatory mechanism that promotes cAMP signalling and hyphal growth post starvation.

**GprH, GprI and GprM influence light-dependent regulation of asexual and sexual development**

The presence of light influences fungal development, where A. nidulans favours asexual conidiation in the light and sexual development in the dark [22]. The A. nidulans wild-type and mutant strains were grown in the presence of light and dark to assess radial growth and asexual conidiation. None of the mutants showed a significant alteration in radial growth (Fig 3A). The wild-type strain produced significantly more asexual conidia in the presence of light than in the dark (Fig 3B). The opposite was true for the ΔgprH mutant, which produced more conidia in the dark and showed reduced conidia production in the light. The ΔgprI and ΔgprM mutants both produced more conidia in the light than dark, but fewer than the wild-type strain. The combination of ΔgprH with ΔgprI, ΔgprM, or both ΔgprIΔgprM, reduced conidiation in the light. However, the combination of ΔgprI and ΔgprM did not reduced conidiation in the light, but instead increased conidiation in the dark.

The production of sexual fruiting bodies, termed cleistothecia, was assessed on sealed minimal media plates incubated in the light or dark (Fig 4A). Under dark conditions inductive of sexual development, ΔgprH increased cleistothecia production (Fig 4B). In the presence of light, ΔgprH and ΔgprI increased cleistothecia production. The combination ΔgprHΔgprI had an additive effect and increased cleistothecia production in the light or dark. However, ΔgprM did not affect sexual development in the light or dark, and when combined with ΔgprH wild-type cleistothecia production was restored. But cleistothecia production remained elevated in the light when the ΔgprM and ΔgprI mutations were combined. Cleistothecia production was maximal in the triple ΔgprHΔgprIΔgprM mutant in the presence of light. Collectively, this again suggests that the GprH signalling mechanism functions in parallel to GprI and GprM, which function in epistasis, to regulate fungal developmental decisions in a light-dependent manner.

**GprH, GprI and GprM mediated, light-dependent, regulation of sexual development during carbon starvation**

Nutrient availability also determines if A. nidulans undergoes either sexual or asexual reproduction [41]. High glucose concentrations promote the production of higher numbers of sexual cleistothecia [41, 42], while light inhibits sexual reproduction [43, 44]. Accordingly, GprH was demonstrated to be a repressor of sexual development during carbon starvation (16), while this starvation condition rescued the developmental defects of the ΔgprD mutant [4]. The atypical production of cleistothecia in carbon starved ΔgprH cultures was found to be dependent on the presence of light. Subsequently, we assessed cleistothecia production in
carbon-starved submerged cultures, exposed to light or dark conditions, for the ΔgprH, ΔgprI and ΔgprM single, double and triple null mutants (Fig 5). Similar to the wild-type strain, neither the ΔgprI nor ΔgprM strains produced cleistothecia under light or dark to assess radial growth and asexual conidiation. Complementation of the single ΔgprH, ΔgprI and ΔgprM mutants with the native gene restored wild-type regulation of sexual development in carbon starved cultures (Fig 5) confirming the phenotype was dependent on the ΔgprH mutation. The combined absence of any two, or all three, of the GPCRs promoted cleistothecia production in carbon-starved light-exposed submerged cultures, even when combining the ΔgprI and ΔgprM mutations, despite the individual ΔgprI or ΔgprM strains failing to produce cleistothecia (Fig 5 and Table 2). This additive effect supports the conclusion that GprI and GprM may function in
Fig 4. GprH, GprI and GprM regulate sexual development on glucose in a light-dependent manner. Conidia of the A. nidulans wild-type (WT), ΔgprH, ΔgprI, ΔgprM, ΔgprHΔgprI, ΔgprHΔgprM, ΔgprIΔgprM and ΔgprHΔgprIΔgprM strains were grown on solid MM and incubated at 37˚C for 24 h in darkness. A) Plates were sealed and transferred into either constant white light or darkness for 96 h to enable cleistothecia (black spherical sexual fruiting bodies) to develop. B) The number of cleistothecia/cm² was determined. Statistical analysis was performed using a two-way ANOVA with Bonferroni post-test when compared to the wild-type strain in each condition (** p < 0.001, * p < 0.01, * p < 0.05).

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Fig 5. GprH, GprI and GprM mediated, light-dependent, regulation of sexual development during carbon starvation. The A. nidulans wild-type (WT), ΔgprH, ΔgprI, ΔgprM, ΔgprHΔgprI, ΔgprHΔgprM, ΔgprIΔgprM and ΔgprHΔgprIΔgprM strains were grown in MM with glucose 2% (w/v) at 37˚C for 24 h. Washed mycelia were transferred to MM containing no carbon source for 8 days, exposed either to white light or darkness. Cleistothecia formation was observed only in submerged cultures exposed to light after 8 days post-carbon starvation, in the ΔgprH, ΔgprHΔgprI, ΔgprHΔgprM, ΔgprIΔgprM and ΔgprHΔgprIΔgprM strains. The repression of cleistothecia production during carbon starvation in the presence of light was restored by complementation of the gprH+ allele. Arrows = cleistothecia magnified in box on top right-hand side.

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Table 2. Cleistothecia production in carbon starved submerged cultures exposed to light or dark. Visual qualitative analysis of the absence (-) or abundance (+) of cleistothecia in the culture media.

| Strain | WT  | ΔgprH | ΔgprI | ΔgprM | ΔgprHΔgprI | ΔgprHΔgprM | ΔgprIΔgprM | ΔgprHΔgprIΔgprM |
|--------|-----|-------|-------|-------|------------|------------|------------|------------------|
| Light  | -   | +++   | -     | -     | +++        | +++        | +++        | +                |
| Dark   | -   | -     | -     | -     | -          | -          | -          | -                |

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GprH represses sexual development by influencing VeA nuclear localization

In *A. nidulans* cleistothecia production predominantly occurs in the dark, while conidiophores are produced in the light [22]. The VeA Velvet protein positively regulates sexual development in a light-dependent manner [26]. In dark, the VeA-VelB complex is imported into the nucleus to activate sexual development. VelB binds to VosA to inhibit asexual development, while the VeA-VelB-LaeA complex activates secondary metabolism [25]. Under light, the transport of VeA to the nucleus is inhibited [17], and as result VelB supports asexual conidiation, while LaeA activity is decreased [24, 45].

The subcellular localization of VeA was evaluated in the wild-type TRMD3.4.17 (*gprH*+) and mutant Δ*gprH* strains, when subjected to carbon starvation under light or dark conditions. The introduction of VeA::GFP into both the TRMD3.4.17 and Δ*gprH* strains did not impact on colony growth or conidiation [17], indicating that the VeA::GFP fusion protein was functional. In the wild-type TRMD3.4.17 strain, VeA::GFP nuclear localization did not increase post carbon starvation under light or dark conditions (Fig 6A and 6B). However, in the Δ*gprH* mutant, VeA::GFP showed increased nuclear accumulation between 30 min and 1 h carbon starvation when exposed to light (Fig 6A and 6B). The nuclear histone h2A of the wild-type and Δ*gprH* VeA::GFP strains were subsequently N-terminal tagged with mRFP (mRFP::h2A). Nuclei were isolated after growth on glucose or during carbon starvation, when exposed to light or dark conditions. Immunoblots confirmed histone isolation and the presence of VeA::GFP in the nucleus. The wild-type strain revealed abundant VeA::GFP nuclear localisation when grown on glucose in the dark, and this was greatly reduced during carbon starvation (Fig 7). The Δ*gprH* mutant showed a lower overall level of VeA::GFP, but a similar regulatory pattern to the wild-type strain when exposed to starvation in the dark. However, in the presence of light, the wild-type strain showed reduced VeA::GFP levels, but an overall similar regulatory pattern of VeA::GFP localisation during carbon starvation. Conversely, the Δ*gprH* mutant showed the opposite regulatory pattern, where VeA::GFP nuclear localisation increased during carbon starvation in the light, correlating with the fluorescence imaging. Therefore, GprH does regulate VeA nuclear localisation and in turn sexual development during carbon starvation in the presence of light.

The allelic veA1 mutation of the veA gene, delays and reduces cleistothecia production [46]. The VeA1 protein lacks the first 36 amino acids at the N-terminus, including the nuclear localization signal, predominantly resides in the cytoplasm [17], and fails to properly regulate sexual development. To examine whether the increased production of cleistothecia observed in the Δ*gprH* mutant was dependent on VeA, a Δ*gprH* strain carrying the veA1 mutation was generated (Fig 8). Cleistothecia production was only observed in carbon-starved cultures exposed to light in the Δ*gprH* mutant carrying the fully functional veA+ allele, and was absent in the Δ*gprH* mutant that possessed the veA1 mutation. Collectively, these data indicate that GprH regulates sexual development via influencing the function of the VeA protein in response to light.

GprH, GprI and GprM have differing effects on ST toxin production

In addition to regulating fungal development, the interaction between LaeA and VeA also coordinates secondary metabolism, including ST and penicillin production, with the presence of light [24, 26, 35, 47]. Among the ST cluster genes, *stcU* is one of the most highly transcribed, and the regulatory *aflR* gene activates the ST gene cluster that leads to the production of ST [31]. In our study, GprH was shown to influence VeA localization in light exposed, carbon-starved cells. Therefore, the role of GprH, as well as that of GprI and GprM, in regulating the
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**A**

| Glucose | Starvation 30 min | Starvation 60 min |
|---------|-------------------|-------------------|
| WT      | ΔgprH             | WT                | ΔgprH          |
| GFP     | HOECHST           | GFP               | HOECHST        |

**B**

| Nuclear accumulation of VeA:GFP Pixel intensity (AU) |
|-------------------------------------------------------|
| Starvation (min)                                      |
| Glu | 30 | 60 | 180 |
| WT  | 1.10 | 1.15 | 1.12 |
| ΔgprH | 1.05 | 1.10 | 1.08 |

**Dark**

**Light**

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The production of ST and penicillin was assessed. In *A. nidulans*, the effect of light on the production of ST depends on the glucose concentration [34].

The epistatic interactions between GprH, GprI, and GprM were examined during incubation under ST-inducing conditions, i.e. static growth on glucose in the dark for 7 days. ST production remained unchanged in the $\Delta gprH$ and $\Delta gprI$ mutants, but was moderately increased in $\Delta gprM$ (Fig 9A and 9B). Combining the $\Delta gprM$ and $\Delta gprH$ mutations had an additive effect and further increased ST production. However, any combination with $\Delta gprI$, including

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**Fig 6. Fluorescence microscopy shows GprH regulates VeA::GFP nuclear localization during carbon starvation in the presence of light.** A) The subcellular localization of the VeA::GFP fusion protein. Conidia of the wild-type TRMD3.4.17 (WT) and the $\Delta gprH$ strains expressing veA::GFP were germinated in MM (glucose 1%), then shifted to MM with no carbon, either constantly exposed to white light or in the dark. Nuclei were stained with Hoechst to verify the VeA::GFP nuclear localization. B) Graphs showing the quantification of the nuclear fluorescence corresponding to VeA::GFP, in the $gprH^+$ and $gprH$ backgrounds. For each condition, 20 nuclei were measured. The error bars indicate standard error. Each experiment was repeated twice. Statistical analysis was performed using a two-way ANOVA with Bonferroni post-test when compared to the wild-type strain in each condition (* $p < 0.01$).

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**Fig 7. Immunoblots confirm GprH regulates VeA::GFP nuclear localization during carbon starvation in the presence of light.** The nuclear histone h2A of the wild-type (WT) and $\Delta gprH$ VeA::GFP strains were N-terminal tagged with mRFP (mRFP::h2A). Nuclei were isolated after growth on glucose or during carbon starvation, when exposed to light or dark conditions. Immunoblots confirmed histone isolation and the presence of VeA::GFP in the nucleus. Coomassie (CB) stained gels show consistent loading of nuclear extracted proteins.

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Fig 8. Fully functional VeA is required for hyperactivation of sexual development in ΔgprH strain. Cleistothecia production was only observed in carbon-starved cultures exposed to light in the ΔgprH mutant carrying the fully functional veA+ allele, and was absent in the ΔgprH mutant that possessed the partially functional veA1 mutation. Cleistothecia were never observed in the wild-type (WT) strain under any carbon starvation conditions. This shows GprH regulates sexual development via influencing the VeA function. Arrow = cleistothecia magnified in box on top right-hand side.

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⊗gprI⊗gprM or ⊗gprH⊗gprI⊗gprM, had the opposite effect and returned ST production to wild-type levels. This suggests that GprM is a major, while GprH is a minor, repressor of ST production, and that the elevated production of ST in their absence was dependent on GprI.

The influence of the three receptor on the transcriptional regulation of known regulators of ST production was assessed after 3 days cultivation under the same ST-inducing conditions. In the ⊗gprM mutant, the expression of the aflR transcription factor was reduced, while the expression of the ST biosynthetic enzyme encoding stcU, and the global laeA regulator, was comparable to the wild-type strain. However, the veA expression was significantly elevated in the ⊗gprI and ⊗gprM mutants (Fig 10). This suggests that the GprI and GprM pathway may influence veA transcriptionally, while the GprH pathway influences VeA nuclear localization and regulation, collectively altering VeA activity which increased ST production in the ΔgprM and ΔgprHΔgprM mutants.

Since GprH, GprI and GprM differentially affected ST production, their influence on the production of another secondary metabolite, penicillin, was assessed using bacterial inhibition
Fig 9. The effect of GPCRs on ST production and expression of ST regulatory genes. A-B) The *A. nidulans* wild-type (WT), ΔgprH, ΔgprI, ΔgprM, ΔgprHΔgprI, ΔgprHΔgprM, ΔgprIΔgprM and ΔgprHΔgprIΔgprM strains were incubated in MM with glucose 1% (w/v) liquid stationary cultures at 37˚C in the dark for 7 days. Presented is the TLC (A) and densitometry (B) analyses of ST production. Statistical analysis was carried out using two-way ANOVA in conjunction with Tukey’s post hoc test (*p* < 0.05). C) The *A. nidulans* wild-type (WT), ΔgprH, ΔgprI, ΔgprM, ΔgprHΔgprI, ΔgprHΔgprM, ΔgprIΔgprM and ΔgprHΔgprIΔgprM strains were assessed for penicillin production. The antibacterial activity of the fungal supernatant was determined by measuring the zones of inhibition *B. calidolactis* C953. No significant difference was observed.

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assays [48]. However, the absence of individual or multiple receptors did not influence the inhibition of bacterial growth and therefore penicillin production (Fig 9C) suggesting that these three GPCRs are dispensable for penicillin biosynthesis in *A. nidulans*.

**Discussion**

Fungi sense extracellular signals to coordinate their development, and cell surface GPCRs coupled with G-protein signalling plays a crucial role in initiating this response [49]. *A. nidulans* possesses 86 putative GPCRs [10], but only a few have been functionally characterised. The *A. nidulans* class III putative carbon receptor, GprD, is structurally related to the *S. cerevisiae* Gpr1 glucose receptor, and functions upstream of cAMP-PKA pathway, regulating conidial germination and hyphal growth on glucose, while repressing sexual development [4, 50]. Gene expression analyses identified additional GPCRs, namely class V gprH and class VII gprM, to be increasingly transcribed during carbon starvation, a condition where GprD is not expressed. GprH was previously shown to be a putative glucose and amino acid receptor functioning upstream of the cAMP-PKA pathway [16, 38]. An additional class V receptor, gprI, remained uncharacterised. This led to the hypothesis that GprH, GprI and GprM may collectively represent a carbon starvation-induced nutrient sensing mechanism. Individually, the
absence of a single receptor caused the over-expression of another receptor, suggesting the existence of redundant or compensatory functions among these carbon starvation-induced receptors. Subsequent genetic analyses demonstrated that GprH functioned in parallel to an epistatic interaction between GprI and GprM, in the regulation of fungal growth, light-dependent sexual development and ST mycotoxin production.

In fungi, GPCRs activate G-protein signalling which initiates the cAMP-PKA pathway [7]. cAMP levels are influenced by glucose availability and increase during glucose starvation. The reintroduction of glucose causes a cAMP burst, which activates PKA [51]. Here, the ΔgprH, ΔgprI and ΔgprM strains did not display increased cAMP levels during carbon starvation, or a burst of cAMP in response to glucose, suggesting that GprH, GprI and GprM all function upstream of the cAMP-PKA pathway. GprH positively influences conidial germination and glucose uptake [16]. Individually, the absence of GprI or GprM had no influence on germination, but the combined absence of both GprI and GprM delayed conidial germination, suggesting epistasis and/or functional redundancy. Intriguingly, the simultaneous absence of all three receptors restored germination defects. The individual ΔgprH, ΔgprI and ΔgprM null mutants all showed a reduction in hyphal growth after the starvation and re-introduction of glucose. In contrast, the combined absence of two or three receptors caused the hyper-activation of hyphal growth during carbon starvation. This correlated with the recovery of cAMP signalling in the double and triple null mutants. Therefore, collectively GprH, GprI and GprM, influence germination and hyphal growth post carbon starvation, but the absence of multiple receptors activates an unknown compensatory cAMP signalling mechanism to promote hyphal growth.

Pheromone sensing, nutrient availability and light determine if A. nidulans undergoes sexual or asexual reproduction [41]. Sexual development requires well-nourished, dark conditions [43]. The GprA/GprB pheromone signalling pathway promotes sexual development [15], while GprD acts as a repressor of sexual development [4], functioning upstream of the pheromone pathway [15]. Similarly, GprH also acts as a repressor of sexual development, but during carbon starvation. The absence of GprH altered the expression of regulators of sexual development, increasing ndsD, and reducing nosA and rosA transcription, resulting in increased cleistothecia production [16]. In contrast to ΔgprH, ΔgprI and ΔgprM mutants did not show enhanced sexual development. However, the combined absence of any two, or all three, receptors resulted in the derepression of sexual development and enhanced cleistothecia production. Again, this provides evidence that GprH functioned in parallel to an epistatic interaction between GprI and GprM in the regulation of sexual development during carbon starvation.

VeA is a light-dependent activator of sexual development [17]. Glucose concentration also affects the light-dependent localization of the VeA transcription factor, as well as the functions of the white collar complex (LreA and LreB) plus the red-light FphA phytochrome [34]. Overexpression of veA causes the increased activation of sexual development [52, 53]. In the dark, VeA accumulates in the nucleus, but this is repressed by the absence of glucose, irrespective of the presence of the GprH receptor. However, during light exposed carbon starvation, the absence of GprH increased the accumulation of VeA in the nucleus, promoting sexual development [24, 26]. This effect on VeA subcellular localization is also reminiscent of that observed in the absence of FphA [18]. The derepression of sexual development caused by the absence of GprD or FphA only occurs in the veA+ background [4, 52]. Introducing the veA1 mutation into the ΔgprH background abolished the derepression of sexual development, confirming that GprH controlled sexual development in a VeA-dependent manner during light exposed carbon starvation. Therefore, both GprH and GprD repress sexual development via regulating VeA function, but under different nutritional states. How these two carbon sensing mechanisms interact with VeA remains unknown.
VeA also interacts with the LaeA methyl transferase that regulates the expression of secondary metabolite gene clusters, while both VeA and LaeA are required for ST mycotoxin production [33, 35]. G-protein signalling similarly affects the ST production [54] and the downstream cAMP-PKA pathway represses the expression of the aflR transcriptional activator of the ST biosynthetic gene cluster, and consequently inhibits ST production [32]. This suggests that GprH, GprI and GprM may influence ST production in a light-dependent manner. The effect of light on fungal ST production also depends on the nutritional state, as by increasing glucose concentration it is possible to bypass the partial repression of ST production caused by light [34]. VeA accumulates in the nucleus independent of light, according to increased glucose availability, and coinciding with an increased production of ST [34]. How the effects of light and glucose availability are integrated in the regulation of ST production remains unknown.

Here, the three receptors differentially influenced ST production, where GprM alone had a major, and GprH a minor, role in the repression of ST production. Interestingly, the increased production of ST in the ΔgprM and ΔgprHΔgprM mutants was dependent on GprI, as the triple ΔgprHΔgprIΔgprM mutant restored wild-type ST production. The transcriptional analysis of genes involved in regulating ST production revealed the GprI and GprM pathway to repress veA expression, which was in contrast to GprH which was shown to regulate VeA nuclear localisation. It remains unknown if GprI and GprM regulate VeA localisation, but this data suggests that these two GPCR mediated pathway may modulate VeA activity via distinct transcriptional and post-translational mechanisms.

In conclusion, important environmental factors such as light and carbon availability affect fungal development and secondary metabolism in A. nidulans. Here, we show how cellular responses to these stimuli occur in an interconnected manner. We revealed GprH, GprI and GprM to be a novel carbon starvation-induced glucose sensing system that functions upstream of the cAMP-PKA, which interacts with the VeA-mediated light sensing mechanism, to regulate fungal development and mycotoxin production. Genetic studies showed the GprH functions in parallel to the epistatic interaction between GprI and GprM in the regulation of sexual development and ST production. Protein-protein interactions studies are now required to dissect how these GPCRs, and their G-proteins [32, 55], interact with the light-sensing systems to control fungal development and ST production. These studies show the complex and integrated structure of fungal mechanisms that perceive different environmental stimuli, through multiple receptors, in turn coordinating fungal development and secondary metabolism, to achieve optimal survival when exposed to environmental versatility. Future studies may reveal how these environment-sensing mechanisms interact to enable fungi to adapt to a fluctuating environment.

Materials and methods

Aspergillus nidulans strains, construction of fungal mutants and growth conditions

Genotypes of the A. nidulans strains used are listed in Supplementary S1 Table and nomenclature of genes is described in [51]. AGB551 and TN02A3 were used as reference strains according to veA+ or veA- backgrounds, respectively. Null gpr strains were generated either by transformation with DNA replacement cassettes [56] or by crossing and selecting recombinants [57]. The gprH, gprI and gprM deletion cassettes were obtained following the fusion-PCR protocols described in [58]. All PCR reactions were performed using TaKaRa Ex Taq DNA Polymerase (ClontechUSA). Supplementary S2 Table lists the specific oligonucleotides used to generate each of the transformation cassettes. Aspergillus fumigatus (Af) pyrG Af and pyrOA Af genes were used as prototrophic selection markers. To study the subcellular
localization of the VeA::GFP fusion protein in the ΔgprH mutant, the TRMD3.4.17 strain containing VeA::GFP [17] was crossed with ΔgprH [16], generating the ΔgprH VeA::GFP strain. The corresponding recombinant strain was selected. To further generate the VeA::GFP mRFP::h2A and the VeA::GFP ΔgprH mRFP::h2A strains, the mRFP::h2A cassette was generated by in vitro recombination as described in [58] with the exception that genes were not replaced by prototrophic markers but were instead N-terminally tagged to mRFP. In this case, the prtA gene was used as a selective marker. Briefly the 5´-UTR and 3´-UTR sequence from histone 2A (h2A) was PCR amplified from genomic DNA from A. nidulans A4 (FGSC). The DNA sequence containing the mRFP plus the h2A gene and its terminator was PCR amplified. The pyrithiamine gene was amplified from pPTRI plasmid (TaKaRa Bio). All fragments were co-transformed, together with plasmid pRS426, previously linearized with EcoRI and BamHI, into S. cerevisiae SC9721 using the lithium acetate method [59]. S. cerevisiae was plated onto selective medium. The whole cassette containing h2A N-tagged to mRFP and the selective marker pyrithiamine was then PCR-amplified from S. cerevisiae genomic DNA, purified and used to transform A. nidulans TRMD3.4.17 or ΔgprH VeA::GFP strains, according to [60]. Transformed candidates were plated in selective medium containing pyrithiamine, purified and checked by both, PCR and microscopy.

To generate the triple null mutant ΔgprHΔgprIΔgprM, the double mutant ΔgprHΔgprI was grown on a plate containing uracil, uridine and the mutagen 5-fluoroorotic acid (5-FOA) (0.75 mg/ml) that allows the recovery of pyrimidine auxotrophs through inactivation of the OMP decarboxylase gene (pyrG) [61]. After three days, sectors showing improved growth were isolated and purified on media containing uracil and uridine. One of the FOA resistant and pyrG- strains was used as recipient of the cassette ΔgprM::pyrG$. For both protoplast mediated transformation and sexual crosses, homologous recombination events were verified by Southern blotting (Supplementary S2 Fig).

To obtain the complemented strains ΔgprH::gprH$ (veA+) and ΔgprI::gprI$ (veA+), the complementing cassettes containing the 5´-UTR region plus the gprH or gprI gene and the 3´-UTR sequences were PCR-amplified from A. nidulans genomic DNA. The auxotopyc marker pyrG was amplified from pCDA21 plasmid and inserted into the complementing cassette as a prototrophic marker. Again, the complementing cassettes were obtained by in vivo recombination in S. cerevisiae. After, the cassettes were fully amplified by PCR and transformed into the ΔgprH or ΔgprI null mutants (both veA$ strains). Positive A. nidulans complementing candidates were selected and purified through three rounds of growth on plates. gDNA was extracted, and the candidates were confirmed by PCR.

Aspergillus complete (YG) and minimal (MM) media, plus the required supplements, were used as described by [62], with or without the addition of 2% agar. Solid YG and MM supplemented for auxotrophies were used to produce conidia for propagation and the maintenance of strains. A. nidulans was grown at 37˚C. Growth tests were performed in MM and colony growth was scored at 37˚C for 48 h (under darkness or constant illumination).

RNA extractions and analysis of gene expression

The expression levels of GPCR genes during carbon starvation was analysed as described by [16]. Briefly, 1x10$^7$ conidia of the corresponding wild-type and mutant strains were inoculated in MM with glucose 1% (w/v) and incubated at 37˚C for 24 h. Next, mycelia were washed and shifted to MM with no carbon source at 37˚C for 4 or 8 h (short-term starvation). Samples at corresponding intervals were frozen in liquid nitrogen and total RNA was extracted using Triazol (Invitrogen) reagent and purified using RNeasy Plant Mini Kit (Quiagen). cDNA was synthesised from 5 μg of RNA using SuperScript III (Invitrogen). Gene expression analysis
was carried out using quantitative real-time PCR (RT-qPCR) according to [63]. cDNA of A. nidulans tubC gene was used for normalization. Corresponding primers for gprH, gprI and gprM are listed in Supplementary S2 Table.

The expression levels of genes involved in ST biosynthesis (aflR, veA, stcU and laeA), was performed as followed. Briefly, 1x10⁶ conidia/mL of the corresponding wild-type and mutant strains were inoculated in MM with glucose 1% (w/v) and incubated at 37˚C, 180rpm, for 72 h under dark condition. Mycelia was filtered and frozen in liquid nitrogen. Total RNA was extracted using Triazol (Invitrogen) reagent and purified using RNeasy Plant Mini Kit (Quia-gen). cDNA was synthesised from 5 μg of RNA using SuperScript III (Invitrogen). Gene expression analysis was carried out using quantitative real-time PCR (RT-qPCR) according to [64]. Primers used for the RT-qPCR analysis are listed in Supplementary S2 Table. The A. nidulans tubC gene was used as a reference.

**cAMP quantification**

1x10⁷ conidia of the AGB551 wild-type and corresponding mutant strains were inoculated in liquid YG media and incubated at 37˚C and 180 rpm for 16 h. Next, mycelia were washed and then incubated in MM lacking a carbon source, at 37˚C for 4 h. After this time, glucose at final concentration of 2% (w/v) was added and incubated for 0.5, 1, 3, 5 or 10 mins. Mycelia were used for cAMP extractions according to (16). The cAMP Biotrak EIA system assays (GE Healthcare) were performed according to manufacturer’s instructions. Total protein content in the corresponding extracts was measured using the Bio-Rad protein assay following manufacturer’s instructions. cAMP concentration was obtained using the ELISA software and data analysis (http://elisaanalysis.com/) and are presented as fmol per μg total protein.

**Microscopy**

To analyse the growth of germlings post-carbon starvation, 1x10⁵ conidia of the A. nidulans strains were incubated in MM with glucose 2% (w/v) for 3 h at 37˚C and then transferred to the modified MM with no carbon and nitrogen sources for 3 h. Next, the starved germlings were shifted to MM for an additional 1 h. Cells were fixed (3% (v/v) formaldehyde and 1.5% (v/v) DMSO in 1x PBS (140 mM NaCl, 2 mM KCl, 10 mM NaHPO₄, 1.8 mM KH₂PO₄, pH 7.4) at room temperature for 10 min and then washed in PBS. For germination assays, 1x10⁵ conidia of the corresponding strains were incubated in MM with glucose 1% for 6 h at 37˚C. After this time, cells were fixed and washed in PBS. All samples were observed using a Zeiss epifluorescence microscope, and the phase contrast bright-field images were captured with AxioCam camera and processed using the AXIOVISION software version 3.1 (Carl Zeiss).

To study the subcellular localization of the VeA::GFP fusion protein, conidia of the wild-type TRMD3.4.17 and the AgprH:VeA::GFP mutant strains were germinated in MM (glucose 1%) and further incubated for 14 h at 25˚C, with white light (28 microEinsteins/m²/s) or in the dark. Next, the cells were shifted to MM with no carbon source for 3 h (light or dark). Samples were observed at 30 min, 1 h and 3 h of carbon starvation. Nuclei were stained with 12 μg/ml Hoechst (Life Technologies, Inc.) to verify the VeA::GFP nuclear localization. DIC images were taken with an exposure of 40 ms, DAPI with 150 ms and GFP with 1000 ms. Images were processed minimally using Wasabi (Hammamatsu Photonics) and ImageJ 1.37v software package.

**Quantification of sexual and asexual development**

To assess cleistothecia production in liquid media, 1x10⁷ conidia of the AGB551, TN02A3 and all the null gpr strains were inoculated in 30 ml of MM with glucose 2% (w/v) and incubated at
37°C for 24 h and 180 rpm. Next, mycelia were filtered and transferred to MM containing no carbon source for 8 days, after which fruiting body production was assessed. Cultures were either exposed to white light (28 microEinsteins/m²/s) or kept in the dark.

Cleistothecia and conidia production was quantified on solid media according to [64] with some modifications. 100 μl of a 1x10⁶ conidia/ml stock were plated MM and incubated at 37°C for 24 h in darkness. Next, one half of the plates was transferred to constant white light for 48 h (asexual) or 96 h (sexual), and the other one was kept in darkness for the same time periods. Plates assessing sexual development quantification were sealed after first 24 h incubation in darkness. To determine the amount of cleistothecia, a defined area of 3.14 mm² was isolated and cleistothecia counted. For the quantification of conidia spores, they were collected in a 2% Tween20 solution and conidia/cm² determined using a Neubauer chamber.

Sterigmatocystin (ST) analysis by thin layer chromatography (TLC)
TLC analyses were carried using procedures previously utilized in [65] with minor modifications. Briefly, A. nidulans strains were inoculated (1x10⁶ conidia/ml) into 25 ml of MM with glucose 1% (w/v) liquid stationary cultures and incubated at 37°C in the dark for 7 days. ST was extracted with chloroform using a 1:1 ratio. Extracts were dried overnight and resuspended in 250 μl of chloroform. Samples were fractionated by TLC on Silica Pre-Coated Polygram Sil G/UV 254 TLC plates (Macherey-Nagal using benzene and glacial acetic acid [95:5 (v/v)] or toluene:ethyl acetate:glacial acetic acid [80:10:10 (v/v/v)] as solvent systems. Plates were sprayed with aluminium chloride (15% in ethanol) and baked at 80°C for 10 min. Metabolites present on TLC plates were visualized under 375 nm UV light. Commercial ST (Sigma-Aldrich) was used as standard. Densitometry of TLC bands was carried out using Gelquant.NET software.

Penicillin (PN) bioassay
The PN bioassay was performed as previously described [48] with some modifications, using Bacillus calidolactis C953 as the test organism. Briefly, A. nidulans strains were inoculated (1x10⁶ conidia/ml) in 20 ml of a seed culture medium, and incubated at 200 rpm for 48 h. After incubation, mycelia were collected using sterile Miracloth (EDM Millipore-Calbiochem) and 1 g of mycelium from each fungal strain was inoculated in 20 ml of fermentation medium. The cultures were incubated at 26°C for 96 h at 250 rpm. After incubation, the cultures were filtered through Miracloth. 300 ml of tryptone-soy agar was supplemented with 50 ml of B. calidolactis C953 culture and plated on five 150-mm-diameter Petri dishes. 10 μl of each culture supernatant was added to 7-mm-diameter wells. The plates were incubated at 55°C for 16 h, and the zones of inhibition (halos) were measured. To confirm that the observed antibacterial activity was due to the presence of PN and not to the presence of other fungal compounds in the supernatant, controls containing commercial penicillinase from Bacillus cereus (Sigma-Aldrich) were used.

Nuclei protein extraction and Western blot analysis
Nuclear proteins were obtained according to Palmer and colleagues [66] with modifications. Briefly, 1 x 10⁶ conidia/ml of the h2A::mRFP;VeA::GFP (WT) or ΔgprH;h2A::mRFP;VeA::GFP strains were inoculated in 500 ml of liquid MM and incubated at 37°C for 16 h (with or without light). Mycelia were filtered, washed three times with sterile water and transferred to MM without carbon source for 3h at 37°C (with or without light). Mycelia were filtered, frozen in liquid nitrogen, and ground to powder. Powdered mycelia were suspended in 20 ml nuclei isolation buffer- NIB (1 M sorbitol, 10 mM Tris-HCl, pH 7.5, 0.15 mM spermine, 0.5 mM...
spermidine, 10 mM EDTA, 2.5 mM phenylmethylsulfonyl fluoride, 1mM DTT, 1 pill/10ml EDTA-free complete Mini protease inhibitor [Roche]) on ice. Samples were centrifuged 10min, 1000 x g at 4˚C. Supernatant was filtered through two layers of Miracloth and centrifuged at 10,000 x g for 15 min at 4˚C. The pellet was resuspended in 1.5 ml resuspension buffer- RB (1 M sorbitol, 10 mM Tris-HCl, pH 7.5, 0.15 mM spermine, 0.5 mM spermidine, 1 mM EDTA, 2.5 mM phenylmethylsulfonyl fluoride, 1mM DTT, 1 pill/10 ml EDTA-free complete Mini protease inhibitor [Roche]) and then centrifuged at 12,000 rpm, 15min at 4˚C. Supernatant was removed and crude nuclei were suspended in 0.4 mL of a modified ST buffer (1 M sorbitol, 10 mM Tris-HCl, pH 7.5, 0.5% NP40, 0.1 mM DTT, 1 pill/10 ml EDTA-free complete Mini protease inhibitor [Roche]). Cellular debris were pelleted by centrifugation (4800 rpm, 30s at 4˚C) and the protein content was measured using a Bradford assay (BioRad).

For each nuclear protein sample, a total of 20μl of GFP by Trap Agarose (ChromoTek) was washed three times with cold ST buffer and incubated for 4h with of 500 μg of nuclear protein extract with shaking at 4˚C. Further, the samples were centrifuged at 5000 x g for 30 seconds and the resin was washed twice with cold ST buffer at 4˚C. Dissociation of bound GFP proteins was performed through addition of SDS-sample buffer and boiling at 95˚C for 5min. Samples were loaded on a 10% SDS-PAGE gel and transferred onto a nitrocellulose membrane using the iBlot 2 Dry Blotting System (Life Technologies) according to the manufacturer’s instructions. For mRFP blotting, a total of 20μg of nuclear protein of each sample was loaded on a 10% SDS-PAGE gel and similarly transferred onto a nitrocellulose membrane. After protein transference, all membranes were blocked for 1 h in 5% (w/v) dry skimmed milk in TBS-T [0.14 M NaCl, 0.02 M Tris, and 0.1% (v/v) Tween 20, pH 7.6] at room temperature. Following, membranes were washed with TBS-T and incubated with a 1:1000 dilution of primary antibody anti-GFP (Sant-Cruz Biotechnology, Inc: Sc-9996) or anti-RFP (ABcam: AB65856) overnight at 4˚C. Primary antibodies were detected using a dilution of 1:10,000 of a horseradish peroxidase (HRP)-conjugated second antibody (Cell Signaling Technology) at room temperature for 1h. Membranes were washed three times with TBS-T and revealed using ECL plus Western Blotting Detection Reagents (Amersham). Coomassie stained gels containing 20 μg of nuclear protein confirmed protein loading.

**Supporting information**

S1 Fig. The corrected gprI (AN8348_M) gene model. A) BLAT alignment of the gene AN8348 on the sense strand of chromosome V of the A. nidulans FGSC_A4 genome shows only all other Aspergillus species and closely related fungi to present an alternative gene model on the antisense strand. B) An alignment of AN8348 with the modified gene model AN8348_M showing their differing orientation, overlap, and intro/exon boundaries. C) InterPro analysis of the AN8348_M gene model yields 7-TM domain characteristic of class V fungal GPCRs.

(TIF)

S2 Fig. Southern Blot analysis confirms deletion of the different A. nidulans GPCRs. A) Deletion of gprI in the ΔgprI and ΔgprIΔgprM strains. B) Deletion of gprI in the ΔgprHΔgprI strain. C) Deletion of gprM in the ΔgprM, ΔgprHΔgprM and ΔgprHΔgprIΔgprM strains.

(TIF)

S1 Table. List of A. nidulans strains used in this work.

(DOCX)
S2 Table. List of primers used in this work.

S1 File. All the raw data presented in the Figs 1–10.

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