Myb Transcription Factors and Light Regulate Sporulation in the Oomycete Phytophthora infestans

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Abstract

Life cycle progression in eukaryotic microbes is often influenced by environment. In the oomycete Phytophthora infestans, which causes late blight on potato and tomato, sporangia have been reported to form mostly at night. By growing P. infestans under different light regimes at constant temperature and humidity, we show that light contributes to the natural pattern of sporulation by delaying sporulation until the following dark period. However, illumination does not permanently block sporulation or strongly affect the total number of sporangia that ultimately form. Based on measurements of sporulation-induced genes such as those encoding protein kinase Pks1 and Myb transcription factors Myb2R1 and Myb2R3, it appears that most spore-associated transcripts start to rise four to eight hours before sporangia appear. Their mRNA levels oscillate with the light/dark cycle and increase with the amount of sporangia. An exception to this pattern of expression is Myb2R4, which is induced several hours before the other genes and declines after cultures start to sporulate. Transformants over-expressing Myb2R4 produce twice the number of sporangia and ten-fold higher levels of Myb2R1 mRNA than wild-type, and chromatin immunoprecipitation showed that Myb2R4 binds the Myb2R1 promoter in vivo. Myb2R4 thus appears to be an early regulator of sporulation. We attempted to silence eight Myb genes by DNA-directed RNAi, but succeeded only with Myb2R3, which resulted in suppressed sporulation. Ectopic expression studies of seven Myb genes revealed that over-expression frequently impaired vegetative growth, and in the case of Myb3R6 interfered with sporangia dormancy. We observed that the degree of silencing induced by a hairpin construct was correlated with its copy number, and ectopic expression was often unstable due to epigenetic silencing and transgene excision.

Introduction

Eukaryotic microbes typically occupy environments in which fluctuating extrinsic factors such as temperature, light, humidity, and nutrients influence progression through the life cycle. Reactions to these conditions are mediated by transcription factors (TFs), protein kinases, G protein-coupled receptors, and other sensors, transducers, and effectors [1,2]. Many environmental responses have evolved to enhance fitness. For example, sporulation at high humidity is common in fungi that make spores that must imbibe water to germinate [3,4]. Often sporulation is suppressed in low oxygen environments which may not favor aerial dissemination [5,6]. In some fungi, solar radiation triggers synthesis of UV-protectants such as carotenoids, or inhibits production of spores that lose viability in light [7]. Other fungi and some slime molds sporulate more intensely in light, perhaps since this signals their presence on the surface of a growth substrate, which may aid spore dispersal [8,9].

How life cycles are integrated with environmental cues is now understood in many fungi, particularly in regards to light [1]. In Aspergillus nidulans, for example, light perception by phytochromes induces Bristle (brlA) protein, a TF that regulates conidiation [10,11]. In Neurospora crassa, the activity of WC (white collar) TFs are regulated by a light-sensing chromophore within the WC protein [7]. Other light-sensing proteins include the cryptochrome/photolyase, opsin, and vivid families [1,12]. Each protein family tends to respond to different wavelengths of light, leading to distinct cellular responses.

Frequently occupying similar niches as fungi, and also sensitive to environment, are oomycetes, which are eukaryotic microbes in the Kingdom Stramenopila. A major genus is Phytophthora, which includes many important plant pathogens [13]. These grow by ramifying hyphae within the host and then form sporangia on plant surfaces, which later germinate by either developing zoospores or directly extending germ tubes. High humidity, which is common at night, is necessary for sporulation in most species of Phytophthora and relatives such as downy mildews [14–16]. Light may also influence sporulation, although data are contradictory with both inhibitory and enhancing effects being reported (rev. in [17]). Sporangia germination is temperature-sensitive, with cool conditions favoring zoospore release over the direct germination [18]. Chemical signals influence zoospore swimming, encystment, and the directional growth of germ tubes [19]. A few genes involved in sporulation have been characterized [20,21], but it is unknown whether their expression or activity is regulated by environmental factors, including light.

In this study, we examine the interplay between light and regulators such as Myb TFs in the sporulation of P. infestans, the species responsible for late blight of potato and tomato. Light is shown to suppress sporulation on plants and artificial media, and...
influence the transcription of sporulation-associated TFs in the Myb family. In an earlier report, we identified 16 R2R3 and R1R2R3-type Myb domain TFs from *P. infestans*, and showed that eight are up-regulated during sporulation [22]. Here, the use of light to synchronize sporulation helped to reveal that one Myb gene (*Myb2R4*) is transcribed earlier than other genes induced during spore development. Chromatin immunoprecipitation and ectopic expression studies supported the role of *Myb2R4* as a regulator of sporulation. Ectopic expression and gene silencing also yielded insight into the activities of other Myb TFs, and technical factors that influence transgene expression and gene silencing.

Materials and Methods

Growth and development of *P. infestans*

Most experiments involved isolate 1306, an A1 mating type strain from tomato in California, USA. Cultures were maintained at 18°C on rye-sucrose agar in the dark, and developmental stages isolated as described [23]. In brief, sporangia were extracted from cultures by adding water and rubbing with a glass rod. To obtain nonsporulating mycelia, sporangia were inoculated into clarified broth and hyphae harvested after 72 hr. Zoosporogenesis was initiated by placing sporangia in 10°C water for 30 min, with zoospores being released after an additional 90 min of incubation. Cysts were obtained by adding 0.25 mM CaCl₂ to zoospores and vortexing for 1 min. Germinated cysts were made by incubating cysts for 6 hr in water at 18°C. Directly germinated sporangia were obtained by placing sporangia for 4 hr in clarified rye-sucrose broth at 18°C. Some studies also involved isolate E13a, an A2 mating type strain from potato in Egypt. Student’s t-test was used to assess differences in growth, sporulation, or germination between treatments or strains.

For studies of sporulation with different light regimes, plates were stored upside down in sealed transparent polystyrene containers containing water-saturated towels. A digital thermometer and hygrometer (National Institute of Standards and Technology traceable) indicated that temperature and humidity were constant at 18°C throughout cultures in darkness. Sporangia were placed in the incubator to 17°C for 12 hr each; to maintain 18°C water for 30 sec, 55°C to 60°C (depending on primer) for 30 sec, and 72°C for 30 sec. Melt curves were generated at the end of each run to test the fidelity of amplification. Expression levels were calculated using the ΔΔCt method, using a constitutive gene (ribosomal protein S3a, PITG_11766) as a control [24,25]. Values are from two biological replicates, with three technical replicates each. qPCR was also used to determine transgene copy number, using DNA extracted as described [26].

Transgenic *P. infestans*

Transformations used the protoplast method [27]. Silencing constructs were based in pTOR (Genbank accession EU257520), in which the ham34 promoter was used to drive transcription of a cassette containing about 400 nt of sense sequences, the Stε2 intron, and antisense sequences [28,29]. pTOR also contains the npl7 gene for selection using G418. Transformants were tested for silencing using semi quantitative RT-PCR, and then RT-qPCR, using the primers shown in Table S2. Over-expression constructs were also based in pTOR, using the relevant gene amplified from cDNA using a C-terminal primer that incorporated a FLAG (DYKDDDDK) tag. Purification of transformants typically involved generating and encysting zoospores (which are usually mononucleate; [30]), plating the cysts on 2% water agar, and transferring single cysts to new rye-sucrose plates lacking G418 using a fine needle. Expression of FLAG-tagged protein was detected by western blot analysis of denaturing polyacrylamide gels using anti-FLAG primary and horseradish peroxidase-conjugated secondary antibodies (Sigma), with detection using chemoluminescence (ECL kit, GE Healthcare). RT-qPCR was used to compare expression levels in transformants and wild-type.

Chromatin immunoprecipitation

The method was adapted from Strahl-Bolsinger et al. [31]. Hyphae in 36 ml stationary liquid cultures were mixed with formaldehyde to a final concentration of 1%, incubated for 15 min using a shaker at 50 rpm, and then for another 5 min with 1.8 ml of 2.5 M glycine. Hyphae were recovered by filtration, washed twice with 20 ml phosphate-buffered saline (PBS), frozen in liquid nitrogen, and ground with 0.4 mm glass beads. Samples were suspended in 0.6 ml of lysis buffer (50 mM HEPES 7.5, 140 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.1% sodium deoxycholate, 1 mM PMSF), sonicated four times for 30 sec at power setting 40 of a Fisher Sonic Dismembrator 550, and clarified by centrifugation at 4650 × g at 4°C for 5 min. Immunoprecipitations used either anti-FLAG antibody or control IgG (Sigma). Prior to being added to the chromatin, 2 μg of antibody was mixed with 10 μl of Protein G magnetic beads (Sigma) in PBS for 1 hr at room temperature.
temperature, washed three times in the same buffer, and resuspended in 50 μl of the same. The antibody/bead/chromatin mixture was incubated overnight at 4°C, and then bead-bound material was recovered using a magnet, washed twice in 0.6 ml lysis buffer, once in 0.6 ml of wash buffer (0.1 M Tris 8.0, 0.25 M LiCl, 0.5% sodium deoxycholate, 1 mM EDTA, 1 mM PMSF, 1 μl of protease inhibitor mix from Sigma), and twice in TE (10 mM Tris-1 mM EDTA pH 8.0). Beads were then resuspended in 0.2 ml TE containing 1% sodium dodecyl sulfate and held at 65°C overnight to reverse cross-links. After adding NaCl to 0.15 M, samples were incubated at 42°C for 2 hr with proteinase K (Sigma) at 1 mg/ml, extracted once with 1:1 phenol/chloroform, precipitated with ethanol, and resuspended in 100 μl water. A fraction was resolved by electrophoresis to check the size of the DNA, and 6 μl aliquots were used for qPCR.

Results

Light influences sporulation timing

To confirm prior reports of the effect of light and obtain materials for further experimentation, tomato leaflets incubated under a 12 hr light/12 hr dark cycle were inoculated with strain 1306 of P. infestans under constant temperature (18°C) and humidity (98% relative humidity). Sporulation began near the end of the fourth day and occurred primarily during the dark periods of days four, five, and six; few sporangia formed in the light (Fig. 1A). The predominance of nocturnal sporulation was also observed on rye-sucrose agar, where sporangia first appeared after four days (Fig. 1B). Cultures in continuous darkness also began to sporulate after four days, but without the periodicity of samples exposed to the light/dark cycle. In studies of strain 1306 and a second strain, E13a, no consistent difference was observed in the total amount of sporangia formed by cultures exposed to light/ dark, constant light, or constant darkness regimes for eight days (Fig. 1C). Light therefore appeared to suppress sporulation, but only temporarily.

Besides affecting the timing of sporulation, light influenced the morphology of cultures. As shown in Fig. 1D, cultures exposed to the light/dark cycle showed a banded pattern, with rings formed every 12 hr. The denser rings were formed in the dark phase, and represent zones with more aerial hyphae, upon which sporangia normally form. However, sporangia were present throughout the culture.

Validation of genes marking sporulation-induced transcription

To help study the interplay of light, sporulation, and transcriptional regulation, four genes with potential roles in spore development were selected for analysis. These were reported previously to be sporulation-induced [22,32]. RT-qPCR was used to confirm that conclusion, and measure expression in other life stages (Fig. 2). The gene encoding the Myb TF named in our prior study as Myb2R3 [22], which corresponds to PTIG_06748 in the Broad Institute genome database, produced modest levels of mRNA in nonsporulating cultures, but high amounts in sporangia and subsequent stages such as zoospores and germinating zoospore cysts. Genes encoding Myb TFs Myb2R1 and Myb2R4 (PTIG_01056, PTIG_00755) and protein kinase Pks1 (PTIG_10884; [33]) exhibited very little mRNA in nonsporulating hyphae, large amounts in sporangia, and declining levels after zoospore release.

Genes induced during sporulation in planta show several patterns

Expression of the genes in P. infestans-infected tomato leaflets exposed to a 12 hr light/dark cycle were measured by RT-qPCR (Fig. 3A). Temperature and humidity were constant during this experiment, as in all work in this paper.

Levels of Myb2R1, Myb2R3, Myb2R4, and Pks1 mRNA were low in leaflets prior to sporulation (48 hr timepoint) and increased just before sporangia appeared. However, the genes exhibited varying kinetics. Myb2R1 and Myb2R3 mRNA rose slightly during the 12 hr period of light preceding sporulation (60 hr), and more during the next dark interval, when sporangia first formed (72 hr). Their mRNA levels then oscillated, declining during each subsequent light period (84 and 108 hr) and rising during each dark period (96 and 120 hr). Pks1 mRNA displayed a similar trend, but with a larger progressive increase during each period of sporulation than Myb2R1 and Myb2R3. A distinct pattern was observed for Myb2R4. Its transcripts also began to rise at 60 hr, just prior to sporulation. However, Myb2R4 showed no light/dark periodicity. Myb2R4 mRNA peaked at 72 hours and fell progressively during the rest of the experiment, while mRNAs of the other genes generally increased along with the number of sporangia.

Several genes not associated with sporulation were measured as controls, as shown in Fig. 3A. Act3a (PTIG_14371) and NPP1 (PTIG_16866) have been described as markers for the biotrophic (early) and necrotrophic (late) stages of infection, respectively [34]. Such patterns were also observed here, but without the oscillations seen for the sporulation-associated genes. CRN2 (PTIG_17199, encoding an effector-like protein) and Tef1 (PTIG_09349, encoding elongation factor-1α) have been described as being expressed throughout the life cycle [25,34], and here they also showed little variation. This confirms that the fluctuations in sporulation-associated mRNAs were not artifacts of the gene used to normalize RT-qPCR, which was PTIG_11706 and encoded ribosomal protein S3a.

A study with additional timepoints provided more insight into when transcriptional changes and sporulation occurred in planta (Fig. 3B). Timepoints were analyzed every 4 hr between 80 and 108 hr post-infection, spanning most of two light periods and one dark period. Most sporulation occurred in the last 4 hr of the night phase, between 92 and 96 hr. Myb2R1 and Myb2R3 mRNA showed slight increases at 84 to 88 hr, and then jumped to much higher levels by 92 hr, just before most sporangia appeared. Myb2R4 mRNA in contrast exhibited its major increase by 84 hr, 4 to 8 hr before the other genes. Myb2R4 transcripts also declined faster during the next light period than those of Myb2R1 and Myb2R3.

The experiments shown in Fig. 3 were performed on detached leaflets, in replicated experiments. Similar results were also observed in whole plants, where sporulation also occurred near the end of the dark period (not shown). For example, Myb2R1 and Myb2R3 mRNAs also oscillated with light and sporangia count in the whole-plant experiments. Myb2R1 mRNA also reached its peak earlier than that of Myb2R1 in the whole-plant studies, and showed little diurnal fluctuation.

Influence of light on transcription and sporulation on artificial media

Trends similar to those detected in planta were seen in P. infestans cultured on rye-sucrose agar (Fig. 4A). Myb2R1 and Myb2R3 mRNA began to rise prior to the appearance of sporangia, had a local peak at 96 hr coincident with the first wave of sporulation, dipped slightly during the next light period, and then rose along
with sporation during the next dark period. As was the case in planta, the relative rise in Myb2R4 mRNA by 72 hr on rye-sucrose media exceeded that of Myb2R1 and Myb2R3. Moreover, Myb2R4 mRNA started to fall by 120 hr, when Myb2R1 and Myb2R3 transcripts were rising. Also, Myb2R4 mRNA did not exhibit light/dark oscillation, unlike Myb2R1 and Myb2R3.

Contrasts between Myb2R4 and the other sporation-associated genes were also observed in cultures experiencing continuous darkness (Fig. 4B) or light (Fig. 4C). In both cases, the level of Myb2R1 mRNA rose faster than the transcripts of Myb2R1, Myb2R3, and Pks1. Furthermore, Myb2R4 mRNA started to decline by 120 hr, while mRNAs of the other genes were still rising. During the continuous dark or light regimes, there was no evidence for 12 hr periodicity in these mRNAs or sporulation.

**Myb2R4 binds the Myb2R1 promoter**

The Myb2R1 promoter bears two copies of a motif that matches the consensus binding site of Myb TFs [22]. Since Myb2R4 is induced soon before other sporulation-induced genes such as Myb2R1, we considered whether Myb2R4 protein might bind the Myb2R1 promoter in vivo to activate transcription. Chromatin
immunoprecipitation (ChIP) studies confirmed that this was the case (Fig. 5). To accomplish this, we generated a *P. infestans* transformant expressing FLAG-tagged Myb2R4 driven by the constitutive ham34 promoter (T10; Fig. 5A). Chromatin was precipitated using anti-FLAG antibody or control IgG, and the resulting DNA was subjected to qPCR using promoter-targeted primers. The Myb2R1 promoter signal was 12 times stronger in the anti-FLAG sample compared to the IgG control (Fig. 5B). In contrast, little enrichment was observed in the anti-FLAG sample for promoters of other genes such as PITG_09284, which encodes an actin-like protein, and PITG_18578, which encodes a protein phosphatase.

Gene silencing suggests that Myb2R3 regulates sporulation

Due to the potential role of Myb2R4 in regulating genes early during sporulation, we tested the effect of silencing it and genes encoding other Myb TFs. Gene disruption has not yet proved feasible in *Phytophthora*, but DNA-directed RNAi using hairpin constructs in stable transformants can yield success [28]. This technique was thus applied to eight Myb genes that we described previously in *P. infestans*, including three expressed in nonsporulating hyphae and sporangia (Myb2R5, Myb3R1, Myb3R3) and five induced during sporulation or in spores (Myb2R1, Myb2R3, Myb2R4, Myb3R5, Myb3R7); wild-type expression patterns of genes not shown in Fig. 2 are described in Xiang et al. [22].

We generated an average of 47 transformants for each gene, and used semiquantitative RT-PCR to identify potential knock-downs and RT-qPCR for confirmation. Convincing knock-downs were observed only for Myb2R3, as shown in Fig. 6A for six representative transformants. Three (T2, T5, and T6) have 5–20% levels of Myb2R3 mRNA compared to wild-type.

Since gene silencing in *Phytophthora* is not well-characterized, we also measured copy numbers of the Myb2R3 transgene in those six transformants to help learn why only some hairpin-derived transformants exhibit silencing. There was a strong correlation ($R = 0.68$) between the level of silencing and copy number (Fig. 6A, lower panel). For example, transformants T2 and T5 had the highest copy numbers and greatest reductions in Myb2R3 mRNA. The range in copy numbers is similar to that seen in a prior study in *P. infestans*, which detected 1 to 130 transgene copies per nucleus with a median of 20 [28].

The three strains silenced for Myb2R3 were checked for abnormalities in growth and development. Each exhibited about 2-fold less sporulation than wild type on rye-sucrose media and in planta (Fig. 6B, C). This is consistent with a role of Myb2R3 as a developmental regulator. Unaffected by silencing were radial growth rates on rye-sucrose media (Fig. 6B, lower), lesion expansion in planta, and spore germination (not shown).

**Ectopic expression of some Mybs impairs growth and is unstable**

Since gene silencing did not succeed for most Myb genes, we also attempted over-expression. This was done for ten genes using...
plasmids designed to produce each Myb with a C-terminal FLAG tag, driven by the *ham34* promoter. Since *ham34* is strong and constitutive [35], it was expected that many transformants would express higher levels of these proteins than wild-type. Seven genes were successfully expressed in this manner. These included four sporulation-induced genes (*Myb3R2*, *Myb2R1*, *Myb2R3*, *Myb2R4*), one expressed in most life stages (*Myb3R6*), one transcribed primarily in hyphae and sporangia (*Myb2R5*), and one expressed in germinating sporangia and zoospores (*Myb3R5*).

Strains over-expressing each of the seven genes produced sporangia capable of releasing zoospores and infecting plants (not shown). However, compared to wild-type and empty vector controls, strains over-expressing *Myb2R1*, *Myb2R3*, and *Myb2R4* showed reduced radial growth on rye-sucrose media (Fig. 7A). Further and more quantitative phenotypic analyses were complicated by the instability of transgene expression. Appearing in cultures were faster-growing sectors (e.g. Fig. 7B), in which transgene expression was reduced. Examples involving *Myb2R4* are shown in Fig. 7C; T10 and T11 are transformants that make FLAG-tagged *Myb2R4*, while T10-R and T11-R are derivatives in which transgene expression was reduced and the slow-growth trait lost. Although T10-R and T11-R displayed diminished expression of the transgene, its copy number was unchanged (Fig. 7D).

We also molecularly characterized a transformant exhibiting instability of expression of FLAG-tagged *Myb2R1*. This is illustrated in Fig. 7F, which shows transformant T20 and a
non-expressing revertant, T20-R. Interestingly, transgene copy number was reduced from 18 copies in T20 to two in T20-R. The slow-growth trait also disappeared in the revertant.

Among single-zoospore (single nuclear) derivatives of the unstable transformants, some maintained expression of both the Myb transgene and the nptII marker, which had been used to select the original transformants. Others expressed only nptII, even though both genes had been introduced on the same plasmid. Single-nuclear derivatives that expressed both transgenes sometimes became unstable in later generations. Therefore, the phenomenon can not be generally explained by the initial transformants being heterokaryons of transformed and nontransformed nuclei. Instead, there appears to be selective pressure to inactivate some Myb transgenes, since over-expression can be deleterious. A similar principle may explain why we failed to obtain strains over-expressing Myb2R2, Myb3R4, or Myb3R7, even after screening 75 transformants.

Ectopic expression suggests that Myb2R4 regulates sporulation

Transformants over-expressing Myb2R4 from the ham34 promoter (T10 and T11) showed about two-fold higher rates of sporulation than wild-type controls (Fig. 7E). The connection between ectopic expression and increased sporulation is strengthened by the fact that strains T10-R and T11-R, which lost most transgene expression, reverted to normal sporulation.

A related phenotype was observed in a transformant over-expressing Myb2R1 (Fig. 7F). The over-expressing strain (T20) showed a 20% increase in sporulation relative to wild-type, while a revertant (T20-R) returned to normal. Although conclusions related to Myb2R1 are tempered by the fact that we could identify only a single over-expressing strain, we present that data to contrast two mechanisms of transgene instability: apparent epigenetic suppression in T10-R and T11-R, and transgene excision in T20-R.

Over-expressing Myb2R4 upregulates Myb2R1

To test the idea implied by our ChIP results that Myb2R4 controls the transcription of Myb2R1, we assessed if over-expressing Myb2R4 increased mRNA levels of the latter. In transformant T10, Myb2R1 mRNA was indeed higher than an empty vector strain by about 9-fold (Fig. 8A). Slight increases were observed for several other Myb genes (Myb2R2, Myb2R3, Myb2R5, Myb3R1, Myb3R3, Myb3R6), but this is probably because these genes are sporulation-induced and T10 produces double the normal amount of sporangia. As controls, we observed normal...
mRNA levels for genes that are not sporulation-induced. These were PITG_09190, which encodes a bZIP TF, and PITG_12808, which encodes an amino acid transporter. We confirmed the connection between Myb2R4 over-expression and Myb2R1 stimulation by examining two additional transformants that over-express Myb2R4, T11 and T12, and the T10-R revertant (Fig. 8B). Myb2R1 mRNA levels ranged from 8 to 20-fold higher than normal in the three over-expressing transformants (T10, T11, T12). Furthermore, Myb2R1 transcripts returned to wild-type levels in T10-R. Over-expressing Myb3R6 weakens sporangia dormancy Unlike the Myb genes mentioned earlier, Myb3R6 is transcribed in all developmental stages (Fig. 9A). We analyzed three transformants over-expressing the Myb3R6::FLAG fusion from the ham34 promoter, that made from 5 to 15 times the normal level of Myb3R6 mRNA (Fig. 9B). Each showed an increased propensity towards premature direct germination followed by repeated [serial] sporulation, as illustrated in Fig. 9C. When sporangia are first harvested from a culture plate they normally appear as in Fig. 9C, panel 1: metabolically active and undescicated, but ungerminated. This was not the case for many sporangia from the over-expressing transformants. As shown in Fig. 9C, panels 2 to 6, many sporangia in 8-day cultures had undergone precocious germination and serial [secondary, tertiary, and quaternary] sporulation, as evidenced by the presence of multiple interconnected germ tubes and sporangia. Although involving a different germination pathway, this is similar in concept to the ability of some species of Phytophthora to resporulate after zoospore release [36]. Serial sporulation was observed in 7 to 15% of sporangia harvested from hyphae of transformants over-expressing Myb3R6 (T60, T61, T62; Fig. 9D). The phenomenon does occur in wild-type P. infestans, but only affects <1% of sporangia under standard growth conditions. The same low frequency of abnormal sporangia was seen in transformants made with the empty expression vector or expressing other transgenes such as Myb2R5 (T50, T51, T52; Fig. 9D). The propensity for serial sporulation increased with culture age (Fig. 9E), but was always much higher in the transformants over-expressing Myb3R6.

Discussion
Like many microbes, P. infestans uses environmental cues to optimize asexual spore production. This study has provided more details about the influence of light on the sporulation; identified Myb2R4, Myb2R1, and possibly Myb2R3 as regulators; and used ChIP and over-expression experiments to demonstrate a direct link between Myb2R4 and Myb2R1. Interestingly, all P. infestans genes encoding Myb2 (R2R3 domain) proteins are sporulation-induced, as are some encoding Myb3 (R1R2R3) proteins [22]. We do not claim that Myb proteins exclusively orchestrate transcription during the transition from hyphae to sporangia in P. infestans, but there are precedents across multiple eukaryotic kingdoms for developmental events being regulated by the concerted action of members of the same TF family [37–39]. Multiple regulators must be involved since sporulation involves many sub-stages such as sporangiophore emergence, coordinated nuclear divisions, development of sporangial initials, papilla formation, establishment of dormancy, etc. [15,40]. Other TF families plus post-transcriptional regulators likely also regulate sporulation. Many protein kinases and some phosphatases are known to be induced at this stage, for example [25,41]. Indeed, a recent study in the related species Phytophthora sojae demonstrated...
that a protein kinase influenced the expression of PsMyb1, which is an ortholog of Myb2R3 of P. infestans [42]. While our analysis found that each of three P. infestans transformants silenced for Myb2R3 were reduced in sporulation, the other study examined one P. sojae strain silenced for PsMyb1 and found that it made more sporangia than wild-type. It is unclear if all PsMyb1-silenced strains would also have that trait, but the P. infestans and P. sojae orthologs may operate dissimilarly as the two species sporulate differently. For example, starvation is needed to induce sporulation in axenic cultures of P. sojae, but not in P. infestans. Also, the sporangia of P. infestans are deciduous, i.e. freely separable from sporangiophores, unlike those of P. sojae [13]. There are also differences in gene expression patterns: while Myb2R3 is sporulation-induced, levels of PsMyb1 mRNA are not much higher in sporulated than nonsporulating cultures [42]. Moreover, while Myb2R3 transcripts remained abundant in P. infestans zoospores, PsMyb1 mRNA was reported to be very low in P. sojae zoospores.

Prior researchers noted that P. infestans produces most sporangia at night, which was attributed to its high relative humidity [14,15]. Nocturnal sporulation presumably benefits P. infestans since its sporangia lack pigments for blocking ultraviolet light and may be more prone to mid-day desiccation [43]. After dawn, fluctuations in temperature and humidity occur that help detach sporangia from sporangiophores, and water films conducive to zoosporogenesis are likely to be present [44,45]. Light maximizes night sporulation by inhibiting daytime development, but humidity seems more critical; once a culture achieves “sporulation competence”, development proceeds only if humidity is above 85–90% [14,46]. Unlike low humidity, light only temporarily delays sporulation, which will occur in cultures receiving continuous illumination. It should be noted that our studies do not address whether light entrains a circadian clock.

Past reports of the effect of light in Phytophthora are conflicting, with light said to inhibit, stimulate, or have no influence on asexual sporulation (rev. in [17]). While some disparity in the literature might be explained by procedural issues, it is also possible that different species may have evolved to respond in varying ways. As noted above for P. infestans and P. sojae, not all members of the genus sporulate in same manner. P. infestans is largely a foliar pathogen and thus may benefit from nocturnal sporulation, but sporulation in the light may promote the aerial dissemination of root-infecting species. Light is known to regulate the balance between asexual and sexual spore formation in some homothallic Phytophthora [47]. In P. infestans, which is heterothallic, we are aware of three controlled studies of the effect of light on asexual sporulation, although none used day/night cycles as employed in our study. These reported that 10 min of light stimulated sporulation [48], continuous daylight reduced sporulation [49], and inhibition resulted from 24 hr of continuous blue light, but not green or red [50].

That blue is the bioactive wavelength may help indicate what P. infestans molecule is the receptor. Of the six types of known photoreceptive proteins [12], only the blue light receptors known as cryptochromes are predicted in P. infestans. Cryptochromes belong to the photolyase/cryptochrome family of flavoproteins and are found in archaea, eubacteria, and eukaryotes [51]. The family is classified phylogenetically and biochemically into cyclobutane pyrimidine dimer (CPD) photolyase, plant CRY, animal CRY, and CRY-DASH (Drosophila, Arabidopsis, Synechocystis, Human) groups [52]. The three putative P. infestans cryptochromes are encoded by genes PITG_01718, PITG_16100, and PITG_16104, and cluster in phylogenetic analyses with animal cryptochromes (Figs. S1, S2).

Light served a useful technical role in this study by facilitating the detection of differences in expression between sporulation-associated genes. Multiple stages of sporulation can be discerned in P. infestans by microscopy, including the emergence of sporangiophore initials, nuclear migration, sporangiophore elongation, nuclear division, swelling of sporangial initials, cytokinesis, and formation of the terminal papilla and basal septum [15,40]. However, these are asynchronous under traditional unilluminated culture conditions, which makes dissecting the stages by RNA analysis impractical. In our light-regulated timecourses, development was spread over longer periods and more synchronous, which helped make it apparent that Myb2R4 was induced earlier during sporulation than Myb2R1, Myb2R3, or Pks1.

The role of Myb2R4 as a regulator of sporulation was supported by our observation that in vivo it bound the promoter of Myb2R1.
Myb2R4 over-expression also stimulated sporulation along with Myb2R1 transcription. Demonstrating Myb2R4 function by stable gene silencing proved elusive, however. We also attempted a transient gene silencing method reported by another group, but without success [53]. Even the use of over-expression to discern Myb2R4 function was challenged by epigenetic events that impaired transgene expression. Loss of expression has also been reported for other transgenes in P. infestans [54], but appeared more problematic for Myb genes, perhaps since over-expression often negatively impacted fitness. With Myb2R4 and Myb2R1, we nevertheless used the phenomenon to our advantage by correlating loss of transgene expression with reversion of increased sporulation phenotypes. It could be argued that over-expression indirectly stimulated sporulation, as a side-effect of reduced vegetative growth.

We observed that inverted repeat transformants with higher copy number were more prone to trigger gene silencing. Similar findings are reported in mammals and plants [55,56]. High copy numbers are common in P. infestans transformants, as most contain tandem repeats of the transforming plasmids [57]. This may also help explain the reduction of transgene copies in some over-expressing strains, since recombination within the repeats could cause excision [58,59]. This is distinct from losses of transforming DNA reported in the related species Phytophthora parasitica, where transgenes persisting as extrachromosomal units were not passed to all daughter nuclei [60]. Nevertheless, vigilance is required.

**Figure 9. Serial sporulation induced by over-expression of Myb3R6.**

A, mRNA levels of Myb3R6 in wild-type, using the same developmental stages as in Fig. 2. Error bars represent standard deviation from three replicates. B, mRNA levels in three over-expressing transformants (T60, T61, T62) and wild-type (WT). C, inferred stages of aberrant germination observed in sporangia harvesting from rye-sucrose agar cultures. Images show sporangium of normal appearance (panel 1), sporangium extending germ tube (panel 2), secondary sporulation (panels 3,4), and formation of a third (panel 5) and fourth sporangium (panel 6). D, Percent of sporangia exhibiting serial sporulation from 8-day cultures of strains expressing Myb3R6::FLAG (T60, T61, T62), a non-expressing revertant of T61 (T61-R), an empty vector transformant (EV), and strains expressing Myb2R5::FLAG (T50, T51, T52). E, Age-dependence of serial sporulation. Data are from a Myb3R6::FLAG transformant, a Myb2R5::FLAG transformant, and an empty vector transformant.

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when testing transgenes in Phytophthora, particularly if a deleterious phenotype is conferred.

Supporting Information

Figure S1 Phylogenetic tree showing relationship between P. infestans cryptochromes and plant CRY, animal CRY, CRY-DASH, and CPD type 1 photolyase groups. The latter four groups are defined as in Daiyasu et al. [52], including the presence of some fungal sequences in the animal CRY group. Also shown are one of the three Phytophthora sojae orthologs and one of two orthologs from Alt homo labachii, which are also oomycetes. Alignments were performed using MUSCLE and a PhyML tree developed using the SEAVIEW program.

Figure S2 Structures of predicted cytochromes from P. infestans.

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Table S1 Corrected gene models. (PDF)

Table S2 Primers used for PCR. (PDF)

Table S3 Numerical values of RT-qPCR data shown in Figs. 3 and 4. (PDF)

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Author Contributions

Conceived and designed the experiments: QX HSJ. Performed the experiments: QX. Analyzed the data: QX HSJ. Wrote the paper: QX HSJ.
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