Unique patterns of mating pheromone presence and absence could result in the ambiguous sexual behaviors of Colletotrichum species

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Abstract

Colletotrichum species are known to engage in unique sexual behaviors that differ significantly from the mating strategies of other filamentous ascomycete species. For example, most ascomycete fungi require the expression of both the MAT1-1-1 and MAT1-2-1 genes to induce sexual reproduction. In contrast, all isolates of Colletotrichum harbor only the MAT1-2-1 gene and yet, are capable of recognizing suitable mating partners and producing sexual progeny. The molecular mechanisms contributing to mating types and behaviors in Colletotrichum are, however, unknown. A comparative genomics approach analyzing 35 genomes, representing 31 Colletotrichum species and two Verticillium species, was used to elucidate a putative molecular mechanism underlying the unique sexual behaviors observed in Colletotrichum species. The existence of only the MAT1-2 idiomorph was confirmed across all species included in this study. Comparisons of the loci harboring the two mating pheromones and their cognate receptors revealed interesting patterns of gene presence and absence. The results showed that these genes have been lost multiple, independent times over the evolutionary history of this genus. These losses indicate that the pheromone pathway no longer plays an active role in mating type determination, suggesting an undiscovered mechanism by which mating partner recognition is controlled in these species. This further suggests that there has been a redirection of the underlying genetic mechanisms that regulate sexual development in Colletotrichum species. This research thus provides a foundation from which further interrogation of this topic can take place.

Keywords: Colletotrichum; sexual reproduction; filamentous fungi; mating pheromones; cognate receptors; gene loss; ancestral state reconstruction

Introduction

The genetic factors that control sexual reproduction in filamentous ascomycetes are largely conserved. This conservation is particularly notable when considering the mating-type (MAT1) locus and its associated genes (Kronstad and Staben 1997). Most described species require the expression of genes from both the MAT1-1 and MAT1-2 idiomorphs in order to sexually reproduce. Those that harbor both sets of genes within a single genome are considered homothallic and are self-sterile (Yun et al. 1999). Alternatively, those that possess either MAT1-1-1 or MAT1-2-1 are typically obligate outcrossers that require an opposite mating type partner in order to engage in sexual reproduction (Turgeon and Yoder 2000; Dyer et al. 2016). These species are thus heterothallic and self-sterile.

The genes present at the MAT1 locus are directly responsible for controlling the expression of the mating pheromones in ascomycete fungi (Zhang et al. 1998; Bobrowicz et al. 2002). In the case of most heterothallic species, the MAT1-1-1 gene positively influences the expression of the α-factor pheromone, while the MAT1-2-1 gene is responsible for α-factor pheromone expression (Zhang et al. 1998; Shen et al. 1999; Bobrowicz et al. 2002; Wilson et al. 2018). Thus, despite the presence of both pheromone genes in the genome, they are expressed in a mating type-dependent manner. Consequently, the expression of these pheromones allows for individuals of opposite mating type to recognize one another as such, subsequently enabling physical interaction and mating (Bolker and Kahmann 1993). This means that while mating type is determined genetically by the MAT1 locus, it is determined physiologically by the pheromone that is being expressed.

Species of Colletotrichum do not exhibit either of these two typical sexual strategies and instead display ambiguous sexual behaviors that are difficult to classify. Sexual reproduction has been shown to occur in numerous Colletotrichum species (Vaillancourt et al. 2000; Guerber et al. 2003; Souza et al. 2010; Barimani et al. 2013), all of which harbor only the MAT1-2-1 gene, while a MAT1-1-1 homolog has never been identified (Vaillancourt et al. 2000; Rodriguez-Guerra et al. 2005; Menat et al. 2012). Consequently, all Colletotrichum species that have been studied to date are of a single mating type- MAT1-2. Despite this, species such as Colletotrichum acutatum are obligate outcrossers, transformed in any way, and that the work is properly cited. For commercial re-use, please contact journals.permissions@oup.com.
with heterothallic-like behavior occurring between two isolates of the same mating type (Damm et al. 2012). This has led to the development of various hypotheses to describe the ambiguous mating behaviors observed in Colletotrichum species.

One hypothesis considering mating strategies in Colletotrichum has been derived from studies on species including Colletotrichum gloeosporioides and Colletotrichum fructicola, and describes a unique form of heterothallism (Menat et al. 2012; Liang et al. 2021). This system, which has been referred to as “unbalanced heterothallism,” is thought to have evolved from a previously homothallic ancestor that has accumulated mutations in genes responsible for self-fertility (Wheeler 1954). Consequently, many different mating identities exist and two individuals are sexually compatible if they harbor complementary mutations. Another hypothesis is that mating identity and cross fertility is determined by two, unlinked loci, rather than the single MAT locus as is seen in most other heterothallic species. Although this has been shown to be true in Colletotrichum graminicola, the actual loci have not yet been identified (Vaillancourt and Hanau 1991; Vaillancourt et al. 2000). In contrast, Colletotrichum lentinus (formerly known as Colletotrichum truncatum or Colletotrichum acetatum) appears to have a classical bipolar mating system that is not linked to the MAT1 locus (Banniza et al. 2018) and has not yet been linked to any other genes.

To further complicate our understanding of the sexual behaviors in Colletotrichum, species such C. gloeosporioides (Weir et al. 2012), C. fructicola (Liang et al. 2021), Colletotrichum fioriniae (Guerber et al. 2003), and C. graminicola (Politis 1975; Vaillancourt and Hanau 1991) have been described as both homothallic and heterothallic. In these species, some isolates are capable of sexual reproduction when cultured alone, while other isolates require a mating partner in order to produce sexual offspring. In C. fructicola, this isolate-dependent behavior is thought to be facilitated by a plus-to-minus switch, with both plus and minus strains being capable of homothallic mating but preferentially undergoing heterothallic mating during periods of hyphal interaction (Liang et al. 2021). However, this system is not well understood, and the presence of homothallic and heterothallic isolates within a single species has made it difficult to elucidate mating type, sexual identity, and mating compatibility in these species. A further complicating factor is that the genus has undergone many taxonomic revisions (Hyde et al. 2009, Marin-Felix et al. 2017; Bhnjun et al. 2021), making it difficult to assign mating strategies to fungi that have been renamed and reclassified many times.

Colletotrichum is divided into 14 currently recognized species complexes (Marin-Felix et al. 2017; Damm et al. 2019; Bhnjun et al. 2021) that accommodate around 220 species (O’Connell et al. 2012; Marin-Felix et al. 2017; Damm et al. 2019). The genus includes important plant pathogens, some of which are recognized as among the top 10 fungal plant pathogens worldwide (Dean et al. 2012). Many Colletotrichum species are hemibiotrophic and have complex life cycles, switching from biotrophic to necrotrophic or endophytic states within the same life cycle (De Silva et al. 2017). These fungi cause a disease known as anthracnose, and infect a diverse range of hosts, including grasses such as maize (Sukno et al. 2008) and sorghum (Pande et al. 1991), fruits such as strawberry (Eastburn and Gubler 1990) and citrus (Rhaiem and Taylor 2016), perennial plants such as pyrethrum (Lelwala et al. 2019), as well as flowering plants like the vanilla orchid (Charron et al. 2018). Symptoms of infection are similarly diverse and include necrotic spots on many different parts of the plant, rotting of the crown and stem as well as seedling blight and brown blotching (Marin-Felix et al. 2017). It is thus important to understand the sexual behaviors of Colletotrichum species, the impact and importance that sexual reproduction plays in their lifecycles and the potential effect ascospore production has on their ability to spread and cause disease.

The aim of this study was to characterize the sexual ambiguity exhibited by Colletotrichum species using a comparative genomics approach. We confirmed the absence of the MAT1-1 gene and the presence of the MAT1-2-1 gene at the MAT1 loci of 31 species across this genus, representing many of the recognized species complexes. In addition, we showed that these species exhibit a unique pattern of gene loss associated with genes of the pheromone response pathway, including the pheromone factors and their cognate receptors. We have shown that these genes have been lost independently, multiple times and we suggest that this could represent a redirection of the underlying genetic mechanisms that regulate the unique mating strategies observed in these fungi.

Methods
Genomes used
This study used genomes from species that represent nine of the 14 recognized Colletotrichum species complexes (Table 1). A total of 25 genomes and their CDS annotations were downloaded from the National Centre for Biotechnology Information (NCBI). This assemblage of genomes included 23 Colletotrichum species, and two Verticillium species that were used as outgroup species in the various analyses conducted. An additional 10 Colletotrichum genomes were downloaded from the NCBI and were annotated using the online version of AUGUSTUS (Stanke and Waack 2003), with default settings and Verticillium longisporum gene models. These 10 additional genomes represented only eight species because three different isolates of C. truncatum were included—C. truncatum 1, C. truncatum 2 (formerly Colletotrichum capsici), and C. truncatum 3. Thus, in total, the 35 genomes represented 31 Colletotrichum species and two Verticillium species.

Phylogenetic analyses
The six gene regions generally accepted as barcoding genes for Colletotrichum were used to generate a species tree (Damm et al. 2019). These regions included act, chi-1, GAPDH, his-3, sod-2, and tub (Supplementary Table S1). Where possible, these sequences were retrieved from Genbank. In the species for which these data were not publicly available, the homologous regions were mined from their respective genomes using the local BLASTn function in CLC Main Workbench V12.0.3. These sequences and the alignments are presented in the supplementary data (Supplementary File S1).

The gene regions were aligned using default parameters in MAFFT V7 (Katoh and Standley 2013). The alignments were subsequently subjected to model testing, using MrModelTest2 V2.4 (Nylander 2004). Thereafter, the alignments were concatenated into a single dataset and analyzed as such. Bayesian inference (BI) analyses were conducted using MrBayes V3.2.7 (Ronquist et al. 2012) using the best substitution model identified by MrModelTest2. The analysis was run for 1 million generations (ensuring average standard deviation of split frequencies <0.01), with 10 parallel runs and four chains. Trees were sampled every 100 generations and 25% of the sampled trees were discarded. Posterior probabilities were then calculated from the remaining trees. The relevant data have been deposited into TreeBASE (Accession number: 28260). The tree was edited in FigTree V1.4.4 (Rambaut 2018) and used in the analyses described below.
including ancestral state reconstruction. It was also used to produce Figures 2, 3, and 6.

**Gene identification and annotation**

**MAT genes**

MAT1-1 CDS sequences and their translated protein sequences from a variety of Sordariomycetes were used in a local BLASTn and tBLASTn searches against all 33 Colletotrichum genomes in CLC Main Workbench V8. These included sequences from *Neurospora crassa* (M33876.1), *Trichoderma reesei* (FJ599756.1), *Fusarium graminearum* (AF318048.1), *Verticillium fungicola* (AB258383.1), and *Podospora anserina* (X64194.1).

The complete MAT1-2-1 CDS sequence from *C. lentis* (KX932470.1) was used as a local BLASTn query against all 33 Colletotrichum genomes in CLC Main Workbench V8. This allowed for the identification of the MAT1-2-1 genes as well as the MAT1 loci in these species. Genes flanking the MAT1-2-1 gene were used as BLASTn queries against the NCBI non-redundant nucleotide database. These results were used to determine the span and gene content of the MAT1 locus as well as the genes that flank the locus. Genes present at the MAT1 locus were translated and subjected to functional domain identification using the NCBI’s Conserved Domains Database (Marchler-Bauer et al. 2015).

In an effort to ensure that all of the possible MAT genes were annotated, the entire MAT locus from each of the Colletotrichum and *Verticillium* genomes was also subjected to a systematic, de novo annotation. The region, which included the two flanking genes, *APN2* and *SLA2*, was annotated using Augustus and Fgenesh, using gene models from *V. longisporum* and *Verticillium hemipterigena*, respectively. Default settings were used for both analyses. Manual annotation was conducted in the few cases where a particular gene was not predicted by either program but where significant similarity allowed for manual gene identification.

**Pheromone genes**

For the identification of the α-factor pheromone, the *Colletotrichum higginsianum* cgq-4 mRNA sequence (XM_018306306.1) and its predicted protein sequence were used as queries in local BLASTn and
tBLASTn searches against all 35 genomes. In species where the gene had not been previously annotated, the BLAST result was used to manually annotate the region likely encoding the pheromone.

Where no usable BLAST results were obtained, a microsynteny approach was used to locate the pheromone gene. Multiple genes found in the 5’ and 3’ regions flanking the α-factor pheromone were identified using a web-based tool called tBLASTn. These sequences were then used as queries in local tBLASTn searches against the remaining genomes. Where relevant, the intergenic regions were subjected to manual identification and annotation.

The predicted protein sequences of the α-factor pheromones were analyzed using various protein identification tools, such as ExPASy ProtScale (Gasteiger et al. 2005) and Phobius (Krogh et al. 2007) to confirm that they possessed the two functional domains associated with the α-factor pheromone—N-terminal hydrophobic region as well as a signal peptide. The sequences were also manually searched for the KEX2 cleavage sites (KR) which flank repeats of an 11 aa region that represents the mature pheromone sequence (Martin et al. 2011).

The α-factor pheromone genes were also identified via a microsynteny-based approach, using the N. crassa α-factor pheromone and its flanking genes. The α-factor pheromone gene in N. crassa is flanked by a cyanase gene (XP.011394492.1) and a hypothetical, EBP2 superfamily protein (XP.960774.1). The predicted protein sequences of these two genes were used as queries in local tBLASTn searches against all 35 genomes. As above, the intergenic regions were subjected to manual identification and annotation. All predicted full-length protein sequences were confirmed to have the defining feature of the α-factor pheromone—the C-terminal CaaX domain. This domain is comprised of a cysteine residue, followed by two aliphatic amino acids and an amino acid of any type (Martin et al. 2011).

### Pheromone receptor genes

The α-factor (preB, XM_018304390.1) as well as the α-factor (preA, XM_018297638.1) pheromone receptor mRNA sequences and their corresponding protein sequences from C. higginsianum were used in various local BLASTn and tBLASTn searches against all 35 genomes.

### Ancestral state reconstruction

In order to determine the ancestral state of the presence or absence of the pheromones and their receptor genes, ancestral state reconstruction was conducted using the unordered, parsimony model in Mesquite V3.61 (Maddison and Maddison 2011). Absence (0) or presence (1) of the genes was mapped onto the BI RIPper, a web-based tool that identifies regions in the genome RIP analysis.

All 35 whole genomes were subjected to analysis using The RIPper, a web-based tool that identifies regions in the genome that have been subjected to repeated induced point (RIP) mutations (Van Wyk et al. 2019). Default parameters were used for all analyses, including those for window and slide size, as well as the composite, product, and substrate values. The average genome-wide GC content and RIP percentages as well as the number of Long RIP Affected Regions (LRAR) and their average GC content were calculated per genome.

### Results

#### The MAT1 locus is conserved across the genus

In all of the Colletotrichum species considered, the MAT1 locus was flanked by the DNA lyase (APN2) and cytoskeleton assembly control (SLA2) genes (Figure 1). In all cases, the MAT1 locus harbored the MAT1-2-1 gene and a previously undescribed secondary MAT1-2 gene. The locus orientation was also highly conserved, with the MAT1-2-1 gene found directly downstream of the APN2 gene (±280–1 100 nt). The only exception was Colletotrichum chlorophyti, where the locus is inverted such that the MAT1-2-1 gene was instead found directly downstream from the SLA2 gene (±1 290 nt).

All of the Colletotrichum MAT1-2-1 genes encoded a protein possessing the HMG (high mobility group) box domain, which is the functional domain associated with this protein in all ascomycete fungi. The secondary MAT1-2 gene present at this locus had not been previously characterized or named and did not show any similarity to other previously described MAT1-2-associated genes. It also did not possess a recognizable functional domain. It has thus been provided with the name MAT1-2-15 in accordance with accepted MAT gene nomenclature (Wilken et al. 2017). Consistent with all previously published research, the MAT1-1-1 gene was not identified in any of the Colletotrichum genomes considered in this study; neither at the MAT1 locus nor elsewhere in the genome.

#### The α-pheromone gene has undergone multiple, independent loss events

The α-pheromone was identified in 17 of the 31 Colletotrichum species and both Verticillium species considered in this study (Supplementary Table S2). All of the identified α-pheromone genes encoded proteins that exemplified the expected structure of the ascomycete α-pheromone factor. In this regard, they possessed a hydrophobic region and signal peptide at the N-terminal. Furthermore, they harbored numerous KEX2 processing sites (KR) as well as one or more repeats of the mature, 11 aa pheromone factor (Supplementary Table S2).

The gene encoding the α-pheromone was found at a fairly conserved location within the genomes of these species, although multiple rearrangements had taken place at this locus (Figure 2). The region downstream of the pheromone gene was highly conserved, with a DUF1640 domain-containing protein present in all species considered. Various other genes encoding known, uncharacterized, and hypothetical proteins (Figure 2, Supplementary Table S3) were found at this locus in species that do not harbor a gene encoding for the pheromone.

In contrast to the conserved nature of the downstream region, the genes found upstream of the α-pheromone varied amongst species (Figure 2). An integral membrane protein as well as a major facilitator superfamily transporter were present in all species considered, but their orientation and location differed between the various species complexes. The presence of a FAD binding domain-containing protein was also conserved at this locus, although it had been lost in Colletotrichum orichophilum and in the species accommodated in the acutatum species complex.

The ancestral state reconstruction analyses showed that the α-pheromone was likely present in the ancestor of Colletotrichum and Verticillium (Figure 3). While this reconstruction showed seven loss events, investigation into the genetic imprints left by these losses indicates more independent loss events. Thus, at least 10 genetic events have occurred at the α-pheromone locus during the evolutionary trajectory of Colletotrichum, each of which
resulted in the loss of the α-pheromone (Figure 4). These events included small changes such as point mutations disrupting the start codon or introducing a premature stop codon. In addition, larger translocations, insertions and/or deletions also occurred and resulted in completely different genes being encoded at this locus.

The first predicted loss event occurred at the split between the orbiculare species complex and all other species of Colletotrichum. In all four species considered from this complex, the α-factor pheromone was absent, and instead, two genes encoding hypothetical proteins are present (Figures 2 and 4). Despite this, the locus was not remarkably different in length compared to the other species. There is also no significant similarity between the region encoding the α-pheromone in any of the other Colletotrichum species and those of the orbiculare complex. It is thus not clear how the gene was lost, but it could have been a large rearrangement or translocation coupled with a deletion.

The loss of the α-pheromone gene in Colletotrichum coccodes represented the third loss of this gene during the evolution of Colletotrichum (Figure 3). While the locus was conserved between C. coccodes and species that do harbor the gene, a point mutation has disrupted the start codon of the putative C. coccodes α-factor pheromone gene (Figure 4). The start codon was mutated from ATG to ACG and would likely not support translation of the protein. Furthermore, the region downstream of the disrupted start codon did not code for any recognizable mature repeats, despite harboring a number of potential KEX2 cutting sites. It is thus likely that this gene is undergoing pseudogenization.

A fourth loss event has occurred in Colletotrichum incanum which is accommodated in the spaethianum complex (Figure 3).
The pheromone region (from integral membrane protein to DUF1640 domain-containing protein) was almost 20 kb in *C. inca-num* and only just over 16 kb in *Colletotrichum tofieldiae*. Despite the much smaller locus, an intact *α*-pheromone gene was identified from the *C. tofieldiae* locus. Interestingly, the encoded protein was much smaller than that of the other *Colletotrichum* species, at only 91 aa long (compared to the others 191–335 aa).

Furthermore, this protein harbors only a single mature pheromone repeat, compared to the two to seven repeats in the other species (Supplementary Table S2). It was particularly relevant that the terminal 43 nt at the 3' region of the *C. tofieldiae* *α*-pheromone gene were well conserved in the *C. incanum* (Figure 5). However, the 5' region of this gene was not conserved between the two species and no definable start codon was present in *C. incanum*. It is therefore possible that there has been an insertion or translocation into the *C. incanum* locus that has replaced the 5' region of *α*-pheromone in this species.

The *α*-pheromone gene was also lost by species residing in the graminicola species complex (Figure 3). In these species, an entirely different gene was present at this locus. In both *C. graminicola* and *Colletotrichum sublineola*, a gene encoding a tetratricopeptide (TTP) protein was present, while this locus in *Colletotrichum falcatum* encoded a hypothetical protein (Figure 4). Similar to the loss in the orbiculare complex species, it is probable that a translocation resulted in the deletion of the *α*-pheromone factor gene and the insertion of new sequences.

Further gene loss events occurred within the acutatum species complex (Figures 3 and 4). The first of these events occurred in *Colletotrichum godetiae*, where a single nucleotide polymorphism (SNP) lead to a nonsense mutation. The putative protein has thus been truncated to only 43 aa. The next two loss events occurred in *C. fioriniae* and *C. acutatum* and both involved the disruption of the start codon, which had been mutated to ATA and ACG, respectively. The final loss event left no genetic imprint and was likely a similar event to the loss of the gene in the graminicola and orbiculare species complexes.

The *α*-factor pheromone flanking genes in *Colletotrichum sansevieriae* were identified at the ends of two different contigs that could not be joined. Therefore, an intact pheromone was not detected in this species. However, the genes as well as the existing intergenic regions are highly conserved between *C. sansevieriae* and *Colletotrichum karstii*, suggesting that the pheromone is likely to present.

The *α*-factor pheromone exhibits less extreme patterns of gene loss

The gene encoding the *α*-factor pheromone was identified in a total of 22 *Colletotrichum* species and both *Verticillium* species (Supplementary Table S4) and was found in the same genomic location in all of them. The gene found directly upstream of the pheromone encodes a cyanate hydratase protein (*CHP*), while the gene directly downstream of the pheromone was eukaryotic rRNA processing protein (*EBP2*). In many of the ten species where the *α*-factor pheromone has been lost, these flanking genes occurred next to one another in the same orientation as those that retained the pheromone gene.

The defining features of the *α*-factor pheromone are its short length and the presence of a highly conserved C-terminal CaaX
In the Colletotrichum species, the sequence of this conserved domain was either CVIL or CVVM (Supplementary Table S4). Although the genes all encode short proteins, there was some variation in the size of the α-factor pheromone protein. For example, Colletotrichum musicola and Colletotrichum plurivorum produced the longest proteins at 109 aa, while species belonging to the orbiculare species complex produced a protein of only 35 aa.

A parsimony-based ancestral state reconstruction showed that the α-factor pheromone was present in the ancestor of Colletotrichum and Verticillium, but that it has subsequently been lost as the species of Colletotrichum have evolved (Figure 6). Although the ancestral state reconstruction showed a total of four loss events, the genetic imprints left during these losses suggest that at least eight independent deletion events have occurred. These events are similar to those that occurred for the α-factor pheromone.

The first loss of the α-factor pheromone occurred in Colletotrichum sojae (Figure 7). Although the C. sojae locus showed high levels of similarity to other species residing in the orchidearum species complex, a 7 nt deletion in the coding sequence resulted in a frameshift mutation. The mutation disrupted the CaaX domain and stop codon. Therefore, any protein encoded by this putative pheromone gene would likely not function as required.

The next α-factor pheromone loss occurred in C. truncatum 2 (Figure 6). While the gene encoding the upstream flanking gene, CHP, is present, the downstream flanking gene is unidentifiable in the C. truncatum 2 genome. This locus has thus been entirely disrupted in C. truncatum 2, leading to the loss of the α-factor pheromone as well. Similarly, no gene encoding this pheromone could be found in C. chlorophyti either, representing the third loss of this gene in Colletotrichum (Figure 6). In this case, the up- and downstream flanking genes are conserved, yet the α-factor pheromone could not be identified in this region or elsewhere in the genome.

The loss of the α-factor pheromone gene in C. coccodes represented the fourth loss of this gene in Colletotrichum species (Figures 6 and 7). In this case, a mutation disrupted the start codon of the gene, changing the start codon from ATG to AGG. The remaining gene sequence was similar to that of closely related species but does not encode for a protein with the terminal CaaX domain.

This pheromone was also lost in the acutatum species complex (Figure 6). However, genetic analysis showed that at least four different deletion events have taken place in this species complex (Figure 7). The first of these occurred in C. godetiae and Colletotrichum salicis, where a disruption of a start codon was
responsible for the loss of this gene. In this case, the start codon was mutated from ATG to GCG and has thus undergone at least two mutation events. Interestingly, a gene sequence similar to that of other species’ α-factor pheromone was detectable after the disrupted start codon and a CaaX domain can be identified. Although there was an alternative start codon, it was found half-way through the gene sequence. Furthermore, there was no stop codon encoded after the CaaX domain and thus any potentially translated protein would be very different from a typical α-factor pheromone. These diagrams are not drawn to scale.

The second loss to occur in the acutatum species complex involves the α-factor pheromone gene of C. acutatum (Figure 7). In this case, a point mutation has introduced a premature stop codon into the gene sequence and would prematurely terminate translation of the putative protein before the terminal CaaX domain, producing a protein of only 31 aa. Interestingly, the third loss involved a similar mutation and occurred in many of the other species residing in the acutatum species complex (Figure 7). In these species, the premature stop codon occurred at position 35. Lastly, the gene was lost in Colletotrichum simmondsii due to another disruption of the start codon, which was mutated from ATG to ACG (Figure 7).

The orbiculare species complex should be mentioned although this gene has not been entirely lost in these species. The α-factor pheromone locus in these species has undergone a deletion of 15 bp that corresponds to the first five amino acids of the α-factor pheromone (Figure 7). An alternative start codon was available, and since no functional domain is known from the N-
terminal of this protein and the C-terminal CaaX domain is still intact, this protein was likely to act as a functional pheromone. However, this would need to be confirmed experimentally.

The cognate receptors display similar patterns of presence/absence as their pheromones

In a similar manner to their respective pheromones, genes encoding the pheromone receptors could be identified in the genomes of only a subset of all Colletotrichum species. In almost all cases, the presence (or absence) of a pheromone gene correlated to the presence (or absence) of the gene encoding the pheromone’s cognate receptor. The receptors, therefore, showed similar patterns of gene presence and absence as their pheromones (Figures 3 and 6). However, there were a few exceptions to this pattern.

With regards to the \(\alpha\)-factor pheromone and its cognate receptor (preB), the exceptions included: C. truncatum 1 and C. tofieldiae, where the pheromone was present while the receptor was absent, and C. godetiae, where the pheromone was absent while the receptor was present (Figure 3). When the \(\alpha\)-factor and its cognate receptor (preA) are considered, three exceptions exist. These were found in C. sojae, C. coccodes, and C. truncatum 2 which, while harboring an intact pheromone receptor, do not possess the corresponding pheromone (Figure 6).

In species where the pheromone receptors were absent, these genes had been lost in similar ways to the pheromone genes. This was particularly true in the case of the \(\alpha\)-factor receptor (preB), where disruptions to the start codon and the presence of in-frame, premature stop codons were common across the genus (Figure 8). Interestingly, these genes have also experienced small to large indel events, a few of which resulted in frameshift mutations, which lead to premature stop codons. Similar genetic events could not be identified at the locus harboring the \(\alpha\)-factor receptor (preA), and it is thus likely that these genes were lost via translocation, insertion, and/or deletion events that left no identifiable trace in the genome.

Colletotrichum species show highly varied levels of genome-wide RIP

Almost all of the genomes considered in this study showed some level of RIP (Table 2), which varied greatly from 0.93% (C. nymphaeae) to 56.02% (Colletotrichum trifolii). The only species that displayed no genomic evidence of RIP was C. sublineola. In general, species residing in the same species complexes showed similar RIP profiles.

The overall GC content of the Colletotrichum genomes was fairly consistent, at around 50% in most of the genomes considered. However, in species where there was significant RIP (>30%),
Figure 7 Genetic events that resulted in loss of the α-factor pheromone in Colletotrichum species clades. A premature, in-frame stop codon truncates the protein in C. nymphaeae before the functional CaaX domain, as shown by ▲. Other C. acutatum clade species and C. coccodes have disrupted start codons as indicated by ▲. C. godetiae and C. salicis as well as C. sojae are missing a stop codon, as indicated by ▲. Interestingly, the C. orbiculare clade species have a 15 bp deletion in the 5' region of the gene, abolishing the start codon that is shared by many of the other species. However, an alternative start codon is present and thus a slightly shorter protein harboring the CaaX domain can likely be produced. These diagrams are not drawn to scale.

Figure 8 Genetic events that disrupted the α-factor pheromone receptor (preB) gene. In many of the species in which this gene has been disrupted, a SNP has abolished the start codon, as shown by ▲. In others, small indels resulted in a frameshift mutation that subsequently produce an in-frame stop codon as shown by ▲. The number in the hexagon refers to the amino acid position at which the stop codon occurs. Deletions are shown by ▲ while insertions are shown by ▲. The number inside the circle refers to the number of nucleotides inserted or deleted. These diagrams are not drawn to scale.
the GC content was considerably lower. For example, members of the orbiculare species complex had some of the highest levels of RIP (45–56%) and, correspondingly, the lowest GC levels (35–38%).

Data availability

The accession numbers of the genomes used in this study are detailed in Table 1, while the accession numbers of the DNA sequences used for the phylogenetic analyses are included in Supplementary Table S1. In addition, Supplementary File S1 contains the sequence data used in the phylogenetic analyses done in this study. The tree files from these analyses were also uploaded to TreeBASE and can be found under the accession number: 28260. Supplemental Material available at figshare: https://doi.org/10.25387/g3.14672052.

Discussion

It has long been apparent that the sexual behaviors of Colletotrichum species are unique amongst filamentous ascomycetes (Politis 1975; Vaillancourt et al. 2000; Menat et al. 2012). These species exhibit ambiguous sexual preferences, with individual isolates in some species having different sexual strategies, as is found in C. fructicola, C. gloeosporioides, and C. graminicola (Vaillancourt and Hanau 1991; Vaillancourt et al. 2000; Menat et al. 2012; Liang et al. 2021). In this study, we have shown that genes encoding the mating pheromones and their cognate receptors displayed an interesting pattern of presence and absence. This might explain the ambiguous and unpredictable nature of sexual reproduction in Colletotrichum.

The only mating-type genes identified in the 31 Colletotrichum species considered in this study were associated with the MAT1-2 idiomorph. While the MAT1-2 locus of these species is comparable to that in other ascomycetes, the complete absence of MAT1-1-associated genes in this genus was the first genetic indicator suggesting that sexual reproduction in Colletotrichum is different to that of other filamentous ascomycetes (Vaillancourt et al. 2000; Rodríguez-Guerra et al. 2005; Menat et al. 2012; O’Connell et al. 2012). The absence of any MAT1-1-associated genes suggests that mating type or sexual identity is no longer genetically determined by the MAT1 locus in these species. While the MAT1-2-1 and

Table 2 The RIP profiles and GC content of the genomes used in this study

| Species               | RIP %  | GC content | No of LRAR | GC content of LRARs |
|-----------------------|--------|------------|------------|---------------------|
| Acutatum species complex |       |            |            |                     |
| C. nymphaeae          | 0.93   | 52.75      | 942        | 30.11               |
| C. simmondsii         | 3.97   | 51.73      | 26         | 24.44               |
| C. forriniae          | 1.42   | 52.49      | 11         | 35.53               |
| C. salicis            | 0.99   | 52.78      | 3          | 38.52               |
| C. godetiae           | 2.89   | 52.09      | 61         | 26.57               |
| C. acutatum           | 2.71   | 52.25      | 45         | 25.89               |
| Graminicola species complex |     |            |            |                     |
| C. graminicola        | 18.09  | 48.42      | 545        | 25.15               |
| C. sublineola         | 0      | 53.11      | 0          | 0                   |
| C. falcatus           | 10.01  | 52.78      | 230        | 27.76               |
| Spaethianum species complex |   |            |            |                     |
| C. tofeldiae          | 4.8    | 52.99      | 110        | 28.12               |
| C. incanum            | 4.64   | 51.9       | 117        | 33.34               |
| Destructivum species complex |   |            |            |                     |
| C. higginsianum       | 3.27   | 54.41      | 117        | 25.89               |
| C. shioi              | 35.92  | 46.26      | 284        | 28.8                |
| C. tanaceti           | 24.1   | 49.19      | 674        | 31.1                |
| No assigned complex   |        |            |            |                     |
| C. orchidiophillum    | 6.22   | 51.06      | 92         | 18.58               |
| C. cocodes            | 1.31   | 53.79      | 0          | –                   |
| C. chlorophyti        | 10.85  | 49.8       | 214        | 25.8                |
| Gloeosporioides species complex | |            |            |                     |
| C. fructicola         | 1.37   | 53.2       | 39         | 19.7                |
| C. asiaticum          | 14.32  | 49.37      | 485        | 23.58               |
| C. camelliae          | 10.76  | 50.01      | 361        | 20.9                |
| C. gloeosporioides    | 2.61   | 51.98      | 107        | 19.7                |
| Boninense species complex |     |            |            |                     |
| C. karstii            | 3.75   | 52.69      | 101        | 23.4                |
| C. sanseveriae        | 11.53  | 50.74      | 38         | 34.88               |
| Trunctatum species complex |   |            |            |                     |
| C. truncatum (1)      | 3.9    | 49.73      | 62         | 27.49               |
| C. truncatum (2) (previously C. capsici) | 5.19 | 48.26 | 123 | 26.72 |
| C. truncatum (3)      | 2.85   | 50.12      | 122        | 35.63               |
| Orchidearum species complex | |           |            |                     |
| C. muscicola          | 5.72   | 54.94      | 133        | 34.87               |
| C. plurivorum         | 0.99   | 55.84      | 15         | 24.47               |
| C. sojae              | 1.02   | 55.9       | 14         | 30.3                |
| Orbiculare species complex |   |            |            |                     |
| C. sidae              | 47.37  | 37.96      | 613        | 19.62               |
| C. orbiculare         | 49.53  | 36.47      | 1067       | 17.91               |
| C. trifolii           | 56.02  | 35.24      | 1045       | 20.85               |
| C. spinosum           | 45.32  | 38.7       | 588        | 19.36               |
| Verticillium (Outgroup) |      |            |            |                     |
| V. tricorpus          | 2.12   | 57.24      | 32         | 32.38               |
| V. dahiae             | 2.59   | 55.34      | 24         | 28.9                |
MAT1-2-15 genes probably remain important for regulating sexual development, it is unlikely that they confer mating type as they do in other species.

The loss of the MAT1-1 idiomorph and its associated genes from all species of Colletotrichum does not appear to have affected the sexual capabilities of a number of these species, which continue to exhibit various forms of homothallic and heterothallic behavior. These species can thus be described as unisexual, as only a single mating type is required for sexual development (Roach et al. 2014). The ambiguous sexual behaviors observed in this genus thus might partly be explained by evolution toward unique sexual strategies that better suit these species and the environments in which they occur. Similar transitions have been observed in other species that exhibit unisexuality, which is thought to be a response to environments where locating a suitable mating partner is difficult (Wang and Lin 2011; Wilson et al. 2015).

The sexual ambiguity and unpredictable nature of mating strategies in species of Colletotrichum can also be linked to the patterns of gene loss associated with the pheromone factors and their cognate receptors. Only 14 Colletotrichum species harbored genes encoding both pheromone factors. Of the remaining species, 11 harbored only one of the pheromones and a further 6 possessed neither. This is in stark contrast to many other studied filamentous ascomycete fungi (Zhang et al. 1998; Shen et al. 1999; Pöggeler 2000; Bobrowicz et al. 2002; Turina et al. 2003; Mayrhofer and Pöggeler 2005; Mayrhofer et al. 2006; Lee et al. 2008; Martin et al. 2011; Wilson et al. 2018) and strongly suggests that mating identity and preference is not under the control of the MAT1 locus and the pheromone pathway in these species. In addition, both homothallism and heterothallism have been documented in these species, behaviors that cannot be linked to a specific pheromone absence/presence pattern. This provides further evidence that the presence and absence of MAT genes and the relevant pheromone pathway genes is not predictive of sexual capabilities or behaviors in Colletotrichum species.

An attractive interpretation of the pheromone presence/absence patterns exhibited in Colletotrichum is that these genes may be implicated in the process of unbalanced heterothallism. It might be envisaged that individuals in the populations of each species could harbor one of the two pheromones and that mating identity and compatibility are determined by these loci. However, the results of this study do not support that hypothesis. Individual isolates residing in the gloeosporioides clade and in which unbalanced heterothallism was first described (Wheeler 1954; Liang et al. 2021), harbor both the α- and β-factor pheromones and their receptors. The same is true for a number of other species of Colletotrichum. In addition, there are some species in which neither pheromone was identified. Thus, presence or absence of one or the other pheromone does not explain the sexual dimorphism in Colletotrichum. Furthermore, the presence/absence patterns are highly conserved within clades. It is consequently unlikely that in all these cases, isolates chosen for whole-genome sequencing would coincidentally exhibit the same patterns of absence or presence as one another in each species complex. In addition, a recent study in C. fructicola (Liang et al. 2021) found that sexually compatible isolates express both mating pheromones and exhibit no sequence differences at these loci, thus supporting the results presented here. It is unlikely that the MAT genes or pheromone genes play a role in unbalanced heterothallism and thus the genes that are involved in this process remain to be identified.

The diversity in the mature pheromone peptide sequences of the Colletotrichum α-pheromone factor was particularly notable. In species such as those of Fusarium and Neurospora, the mature pheromone peptide is highly conserved (Martin et al. 2011). This is even true for N. crassa and Sordaria macrospora which, despite being accommodated in different genera, produce identical α-factor pheromones. This is in contrast to the high level of diversity seen in the β-factor pheromones of Colletotrichum species. This is true both of the sequence and number of mature pheromone repeats encoded by the gene. For example, C. higginsianum and Colletotrichum tanaceti reside in the same species complex and yet encode two and three mature repeats respectively and share no mature pheromone repeats. Furthermore, Colletotrichum asiamum produces an α-factor protein harboring a total of six mature repeats, made up of four different amino acid sequences. This suggests that this region is rapidly evolving and may contribute to the unpredictable nature of mating in this genus.

Despite sexual cycles not being apparent in many of the Colletotrichum species considered in this study, RIP profiles suggest that sexual reproduction is more common in the genus than is reflected by known sexual states. RIP is a process that is thought to exclusively occur during the pre-meiotic phase of sexual reproduction in filamentous ascomycete fungi and may thus be predictive of the sexual cycle (Hane et al. 2015). Importantly, RIP may indicate that sexual reproduction has taken place in the past and may consequently not be predictive of sexual recombination actively occurring in an extant population. Some Colletotrichum species, particularly those belonging to the orbiculare species complex, have surprisingly high RIP values, up to 56% in C. trifolii. This is significantly higher than other reports of genome-wide RIP, including those for N. crassa (15%) and various Fusarium species (up to 6.4%) (van Wyk et al. 2019). This is particularly interesting given that the sexual cycle has not been observed in any of the four orbiculare clade representatives included in this study. The significance of these high RIP levels deserves further investigation as it may have relevant implications in the evolutionary history and future of this genus.
The three C. truncatum genomes considered in this study displayed various interesting differences that are worth noting. The C. truncatum 1 and C. truncatum 2 isolates were both isolated from Capsicum annuum, while C. truncatum 3 was isolated from soybean. Despite this, both C. truncatum 1 and 3 harbored both mating pheromones as well as the α-factor receptor, while C. truncatum 2 harbored neither pheromone. In fact, the loci at which the pheromone genes are typically found in other Colletotrichum species were entirely disrupted in C. truncatum 2, with significant rearrangement and gene loss events having taken place. However, when the MAT1 locus is considered, C. truncatum 1 is different from the other two isolates, with a significantly longer locus harboring a 4Kb insertion. This is of significance because C. truncatum 2 was originally described as C. capsici but has since been reduced to synonymy with C. truncatum. These results may provide support for treating C. capsici as a discrete species, and they also provide preliminary evidence of a C. truncatum × C. capsici hybrid.

Conclusions

The ambiguous sexual behaviors exhibited by species of Colletotrichum are potentially the result of a unique pattern of gene presence and absence. Unlike most other studied filamentous ascomycetates, both the pheromones and their receptors are not consistently present in the genomes of species in this genus. This has implications for how these fungi can sexually reproduce, particularly in the manner in which they recognize potential mating partners. Future research should include transcriptomic analyses to determine the expression patterns of the genes that are present, particularly in isolates known to be sexually compatible.

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Conflicts of interest

None declared.

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