Comparative Genomic Analysis of Ochratoxin A Biosynthetic Cluster in Producing Fungi: New Evidence of a Cyclase Gene Involvement

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The widespread use of Next-Generation Sequencing has opened a new era in the study of biological systems by significantly increasing the catalog of fungal genomes sequences and identifying gene clusters for known secondary metabolites as well as novel cryptic ones. However, most of these clusters still need to be examined in detail to completely understand the pathway steps and the regulation of the biosynthesis of metabolites. Genome sequencing approach led to the identification of the biosynthetic genes cluster of ochratoxin A (OTA) in a number of producing fungal species. Ochratoxin A is a potent pentaketide nephrotoxin produced by Aspergillus and Penicillium species and found as widely contaminant in food, beverages and feed. The increasing availability of several new genome sequences of OTA producer species in JGI Mycosmos and/or GenBank databanks led us to analyze and update the gene cluster structure in 19 Aspergillus and 2 Penicillium OTA producing species, resulting in a well conserved organization of OTA core genes among the species. Furthermore, our comparative genome analyses evidenced the presence of an additional gene, previously undescribed, located between the polyketide and non-ribosomal synthase genes in the cluster of all the species analyzed. The presence of a SnoaL cyclase domain in the sequence of this gene supports its putative role in the polyketide cyclization reaction during the initial steps of the OTA biosynthesis pathway. The phylogenetic analysis showed a clustering of OTA SnoaL domains in accordance with the phylogeny of OTA producing species at species and section levels. The characterization of this new OTA gene, its putative role and its expression evidence in three important representative producing species, are reported here for the first time.

Keywords: ochratoxin A genes, Aspergillus, Penicillium, genomic analysis, SnoaL domain, cyclase
INTRODUCTION

Next-generation sequencing (NGS) technologies have opened a new era in the study of biological systems by significantly increasing the catalog of fungal genomes sequences available for analysis. The Fungal Genomics Program\(^1\) of the United States Department of Energy (DOE) Joint Genome Institute (GI) has partnered with the international scientific community to support genomic exploration of the fungal kingdom and to help in addressing this lack of knowledge (Grigoriev et al., 2011). The exponential growth of data resulting from several large-scale genomics initiatives, like the 1000 Fungal Genomes Project\(^2\) and the Aspergillus Whole Genus Sequencing Project (Vesth et al., 2018; Kjaerbølling et al., 2020, 2018), have provided novel genomic information of a vast amount of fungal species. In parallel with increased genome sequencing throughput, transcriptomic data from fungi have become easily attainable with RNA-seq approaches enabling current gene modeling pipelines to combine genome sequences with deep transcript sequencing data. This integrated approach has led to improvements in gene model generation, allowing the identification of previously "unseen" genes that are often comparatively small in size and missed by prior automated annotation pipelines (Grigoriev et al., 2014). This revolutionary approach to genome characterization has enabled a large-scale in silico identification of putative genes encoding novel enzymes, metabolic pathways, and bioactive compounds.

Secondary metabolites (SMs) produced by microorganisms are of major interest because of their suitability for the treatment of infectious diseases, as immunosuppressants, or as sources of new and innovative therapeutic agents; however, they are also implicated in food safety and human health as mycotoxins or bacterial toxins, or antibiotics (Brakhage and Schroeckh, 2011; Frisvad and Larsen, 2015). During the 1990s, a variety of SMs were genetically characterized and the clustered arrangement of genes involved in the biosynthesis of a single SM was studied (Sidhu, 2002). Gene cluster discovery in fungi was complex and time-consuming in the pre-genomic era, since their study involved cumbersome traditional molecular methods. Conversely, in the genomic era, the investigations of genome encoded biosynthetic potential have accelerated and the potential for exploitation of biosynthetic gene clusters gained a great attention.

Comparative analysis of predicted genes has led to the identification of many SM clusters of known metabolites, revealing the genetic basis of their biosynthesis and regulation (Kjaerbølling et al., 2018). On the other hand, the same approach has led to the prediction of novel cryptic clusters for still unknown microbial metabolites, shedding light on new biosynthetic pathways never explored before (Khaldi et al., 2010; Metzker, 2010; Cacho et al., 2014; de Oliveira and Ocaña, 2015). However, most of the clusters identified by genome analysis require more investigation before we completely understand the pathway steps and the regulatory network behind the metabolite biosynthesis (Brakhage, 2013).

An appropriate example for how the genomic approach has led to the identification and characterization of biosynthetic gene clusters (BGC), is ochratoxin A (OTA), primarily identified in Aspergillus niger, and subsequently in A. carbonarius (Pel et al., 2007; Baker et al., 2009). Ochratoxin A is a well-known mycotoxin with wide distribution on food and feed, including cereal products, grapes and by-products, coffee, beverages, cocoa, nuts, dried fruits, and cured meat (Jørgensen, 2005; el Khoury and Atoui, 2010). Ochratoxin A is nephrotoxic, hepatotoxic, embryotoxic, teratogenic, neurotoxic, immunotoxic, genotoxic, and carcinogenic with species and sex-related differences (IARC, 1993; Castegnaro et al., 2006; Malir et al., 2016).

The analysis of A. carbonarius genomic data revealed the key role of three genes (AcOTAPks, AcOTANrps, and AcOTAhal) in the OTA biosynthesis (Gallo et al., 2012, 2017; Ferrara et al., 2016), and the involvement of two genes encoding for a cytochrome P450 oxidase and a bZIP transcription factor has been demonstrated recently in six different OTA producing species (Wang et al., 2018). This evidence, supported by molecular genetic approaches, has defined the role of these five core genes as essential for OTA biosynthesis. Moreover, a comparative analysis of OTA-biosynthetic clusters in A. steynii, A. westerdijkiae, A. niger, A. carbonarius, and P. nordicum has revealed substantial synteny in cluster organization (Gil-Serna et al., 2018).

To date, OTA is known to be produced by several species of genus Aspergillus – more than twenty mainly within Sect. Nigri and Circumdati, and at least three Penicillium species (P. nördicum, P. thyrioida, P. verrucosum). The recent availability in JGI Mycocosm and/or GenBank of several new genome sequences within Aspergillus and Penicillium OTA producer species has encouraged us to extend the analysis of cluster organization to 21 OTA producing species, 19 Aspergillus and 2 Penicillium. The extensive analysis of the genomic region related to OTA cluster genes, indicated presence of an additional previously undescribed gene, with a “SnoA-like cyclase” domain located between the polyketide synthase (PKS) and non-ribosomal peptide synthase (NRPS) genes.

The role of cyclization in polyketide biosynthesis is complex and important. There is limited data regarding the role of specific cyclase genes in secondary metabolite biosynthesis. More often, cyclization is thought to happen either spontaneously via intramolecular interactions within the growing polyketide or via cyclization domains that are part of the PKS; alternative hypothesis could be the involvement of an external cyclase gene respect to the PKS domain structure.

In this respect, the characterization of this SnoA-like cyclase gene, its phylogenetic relationships, putative role and expression evidence in three important representative species under producing conditions are reported for the first time in this manuscript.

\(^1\)http://jgi.doe.gov/fungi

\(^2\)http://1000.fungalgenomes.org
Materials and Methods

Ochratoxin Cluster Analysis in Ochratoxigenic Fungi

The analysis of the complete OTA cluster of producing species was carried out by retrieving sequence information and gene annotation from JGI Genome Portal for A. sclerotiniger CBS115572 v1.0, A. carbonarius ITEM 5010 v3.0, A. niger ATCC 13496 v1.0, A. steynii IBT 23096 v1.0, A. muricatus CBS 112808 v1.0, A. roseoglabulosus CBS 112800 v1.0, A. flocculosus CBS 112785 v1.0, A. pulvericola CBS 137327 v1.0, A. elegans CBS 116.39 v1.0, A. welwitschiae CBS139.54b v1.0, A. affinis CBS 129190 v1.0, A. cretensis CBS 112802 v1.0, A. subramanii CBS 138230 v1.0, A. albertyensis IBT 14317 v1.0, A. alliaceus CBS 536.65 v1.0, A. sulphureus (fresenii) CBS 550.65 v1.0, A. sclerotialcarbonarius CBS 121057. For A. westerdijkiae, and P. nordicum, we considered the gene annotation reported by other authors (Gil-Serna et al., 2018; Wang et al., 2018) and compared it with the data available at JGI for A. westerdijkiae CBS 112803 v1.0 and P. nordicum DAOMC 185683 v1.0. Sequence information for A. ochraceus FC-1 (GCA_00489945.1) and P. verrucosum BFE808 (GCA_000970515.2) were retrieved from NCBI database. For each species the main characteristics of the biosynthetic genes were described including the information related to DNA, mRNA and proteins. Gene features and annotations were retrieved by BlastX and manually cured, when not available.

Fungal Strains and Growth Conditions

Fungal strains used were A. carbonarius ITEM 5010, A. westerdijkiae ITEM 9607 and P. nordicum ITEM 9634 from the Agro-Food Microbial Culture Collection of the Institute of Sciences of Food Production, CNR, Bari, Italy. Fungal strains were routinely grown on PDA (Oxoid, United Kingdom).

For confirmation of PKS cyclase gene (otaY) expression under OTA permissive condition, 100 µl of a conidial suspension (10^6 conidia/ml) of each strain were inoculated on yeast extract sucrose (YES) agar plates (yeast extract 20 g/l, sucrose 150 g/l, agar 20 g/l) overlaid with sterile cellophane membranes. Incubation was carried out in the dark at 25°C. Fungal mycelium was harvested after 5 days post inoculation (dpi), then frozen in liquid nitrogen and stored at −80°C for RNA extraction. Triplicate cultures were prepared and analyzed for each strain.

Nucleic Acid Extraction, cDNA Synthesis, RT-PCR and Sequencing

Genomic DNA (gDNA) was extracted using the Purelink Genomic Plant DNA kit (Invitrogen, San Diego, CA). Total RNA was extracted from frozen mycelium ground in liquid nitrogen using the RNEasy kit (Qiagen, Hilden, DE) according to the manufacturer's protocol. Any possible DNA contamination was removed by RNase-free DNase I (Qiagen) digestion. RNA aliquots were stored at −80°C. First-strand cDNA was synthesized using 1 µg of total RNA with SuperScript IV First-Strand Synthesis System (Invitrogen, San Diego, CA) according to the manufacturer’s protocol. Reverse transcription-PCR (RT-PCR) was used to analyze the expression of otaY transcripts in A. carbonarius, A. westerdijkiae and P. nordicum strains. Amplifications were carried out on gDNA and cDNA using the Platinum SuperFi II PCR Master Mix (Invitrogen, San Diego, CA). The primer pairs listed in Table 1 were used under the following conditions: 98°C for 30 s, followed by 35 cycles of 98°C for 10 s, 60°C for 10 s, and 72°C for 30 s, followed by a final extension step at 72°C for 5 min. Amplicons were checked by 1.5% (w/v) agarose gel electrophoresis and sequenced using the BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystem, United States).

The deduced amino acid sequences were determined using the EMBoss Sixpack tool. Protein sequences were aligned by using muscle algorithm and the functional domain was detected by SMART analysis.

Comparative Genome Analysis for Cyclases in Fungi

During the analysis of genomes sequenced as part of the Whole Aspergillus Genus Sequencing Project, gene clusters predicted to encode the enzymes involved in OTA biosynthesis were identified in several of the genomes. Within those clusters, we noted the conserved presence of a putative SnoaL domain containing cyclase. In some of the previously published Aspergillus genomes, the cyclase gene was present, but unidentified by prior versions of the gene modeling software. In these cases, the cyclase gene was manually curated. Following the discovery of the SnoaL domain containing cyclase we searched all available Aspergillus genomes for SnoaL cyclases and downloaded all those in the range of 160–230 amino acids to perform phylogenetic analysis.

Multiple Sequence Alignment and Phylogenetic Analysis of Cyclase Domain

A total of 863 amino acidic sequences of SnoaL domains were retrieved from Aspergillus fungal genomes portal (MycoCosm, DOE-JGI) and subsequently aligned using the MAFFT software Version 7 (Katoh and Standley, 2013). The phylogenetic analysis of all the SnoaL domains was conducted by the Maximum Likelihood method using EMBOSS Sixpack tool6.

Table 1: Primer sequences for RT-PCR.

| Primer name       | Nucleotide sequence                  |
|-------------------|--------------------------------------|
| Aspca_cycl_F      | 5′-ACCATCCTCACCACCGTTGTG-3′          |
| Aspca_cycl_R      | 5′-CGGTATCTCTCAACCCACCAAA-3′         |
| Aspwt_cycl_F      | 5′-TGATCTCAAGTACCGGCT-3′             |
| Aspwt_cycl_R      | 5′-CTCCCTCTTCATCTCGCACT-3′           |
| Pnord_cycl_F      | 5′-TCCGCTGCTTGAACTGCTTCT-3′         |
| Pnord_cycl_R      | 5′-TCATTGCTCCTCGGATGATG-3′           |

6https://www.ebi.ac.uk/Tools/st/emboss_sixpack/  
4http://smart.embl-heidelberg.de/
Likelihood (ML) using RaxML hybrid code (Stamatakis, 2014) at the CIPRES Science Gateway (Miller et al., 2010). Bootstrapping was performed to assess the robustness of the phylogeny. The most robust approach to infer phylogenetic relationship was ML analysis by using the RAxML algorithm, setting the bootstrap analysis to 1000 runs, the GAMMA Model parameters were used up to an accuracy of 0.1000000000 Log Likelihood units.

For our OTA-specific phylogenetic study, we selected the 23 OTA clustered SnoaL cyclases out of a total of 863 from our preliminary analysis (Supplementary Figure 1). We also included two SnoaL cyclases from non-OTA strains of A. parvulus, A. cervinus. Finally, we included a SnoaL cyclase from Metarhizium robertsi found by genomic analysis to belong to a putative-partial OTA-like cluster. For this analysis the aa sequences domains were aligned both with Clustal and MUSCLE alignment in MegaX software (Kumar et al., 2018) by trimming manually the external not well aligned part of the obtained alignment; the validity and the robustness of the alignment was checked on T-Coffee server2. The best substitution model was calculated in Mega X. The evolutionary history was inferred by using the Maximum Likelihood method and JTT matrix-based model (Jones et al., 1992) by the use of Gamma distribution and Invariant sites. There were a total of 115 positions in the final dataset. Evolutionary analyses were conducted in MEGA X. Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using a JTT model, and then selecting the topology with superior log likelihood value. A discrete Gamma distribution was used to model evolutionary rate differences among sites [5 categories (+G, parameter = 1.3800)]. The rate variation model allowed for some sites to be evolutionarily invariable ([+I], 12.17% sites). The same set of alignments was converted in nexus format and run on MrBayes software (Ronquist et al., 2012) to conduct the Bayesian Posterior Probability analysis for the evolutionary tree. The Dayhoff substitution rate was set with 4 categories of Gamma distribution with invariable sites uniform distributed, the tree. The Dayhoff substitution rate was set with 4 categories of Gamma distribution with invariable sites uniform distributed, the number of sequences were 26 and 126 characters with 113 unique site patterns. This analysis involved 26 amino acid sequences. All positions containing gaps and missing data were eliminated (complete deletion option).

RESULTS

Annotation of OTA Cluster Structure in Ochratoxigenic Species

Analysis of the OTA cluster was performed on 21 ochratoxigenic species, most of them belonging to the Aspergillus genus, with only two species belonging to Penicillium genus (Table 2). The length of each OTA core cluster in the species analyzed varies from 19 to 24 kbp, with differences due to variance in the intergenic space and gene model sequences. The core OTA biosynthetic cluster is composed of five genes, highly conserved in all the ochratoxigenic species. The genes encode a PKS,

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2http://tcoffee.crg.cat/apps/tcoffee/index.html

### Table 2 | List of main features of the genes in OTA cluster of the 21 species analyzed.

| Species (core cluster length) | Gene | DNA length(bp) | Number of exons | cDNA length(bp) | aa |
|---|---|---|---|---|---|
| Aspergillus sclerotiorum CBS115572 | otaA | 8012 | 9 | 7614 | 2537 |
| (21156 bp) | otaY | 420 | 1 | 420 | 139 |
| Aspergillus carbonarius ITEM 5010 | otaA | 8063 | 9 | 7626 | 2541 |
| (21508 bp) | otaB | 5638 | 2 | 5586 | 1861 |
| (23497 bp) | otaC | 1763 | 6 | 1542 | 513 |
| Aspergillus niger ATCC 13496 | otaA | 8173 | 9 | 7665 | 2554 |
| (21670 bp) | otaB | 5692 | 2 | 5643 | 1880 |
| (23434 bp) | otaC | 1809 | 6 | 1551 | 516 |
| Aspergillus steynii IBT 23096 | otaA | 8048 | 9 | 7611 | 2526 |
| (23671 bp) | otaB | 5689 | 2 | 5640 | 1879 |
| (23599 bp) | otaC | 1822 | 6 | 1542 | 513 |
| Aspergillus muricatus CBS 112808 | otaA | 8083 | 9 | 7647 | 2548 |
| (23696 bp) | otaB | 5694 | 2 | 5646 | 1881 |
| (21508 bp) | otaC | 1822 | 6 | 1542 | 513 |
| Aspergillus roseoalbatus CBS 112800 | otaA | 8058 | 9 | 7653 | 2550 |
| (23599 bp) | otaB | 5699 | 2 | 5652 | 1883 |
| (21671 bp) | otaC | 1838 | 6 | 1542 | 513 |
| Aspergillus flocculosus CBS 112785 | otaA | 8032 | 9 | 7602 | 2533 |
| (23434 bp) | otaB | 5689 | 2 | 5640 | 1879 |
| (23410 bp) | otaC | 1825 | 6 | 1545 | 514 |
| Aspergillus niger CBS 137327 | otaA | 8080 | 9 | 7650 | 2549 |
| (23410 bp) | otaB | 5689 | 2 | 5640 | 1879 |
| (23410 bp) | otaC | 1824 | 6 | 1545 | 514 |
| Aspergillus westerdijkiae CECT 2948 | otaA | 8095 | 9 | 7608 | 2535 |
| (24204 bp) | otaB | 5701 | 2 | 5652 | 1883 |

(Continued)
| Species (core cluster length) | Gene    | DNA length (bp) | Number of exons | cDNA length (bp) | aa |
|-----------------------------|---------|-----------------|-----------------|-----------------|----|
| Aspergillus ochraceus FC-1  | otaC    | 1821            | 6               | 1545            | 514|
|                             | otaA    | 8143            | 9               | 7656            | 2551|
| (24204 bp)                  | otaC    | 1827            | 6               | 1545            | 514|
|                             | otaD    | 1776            | 2               | 1479            | 492|
| Aspergillus elegans CBS 116.39 v1.0 | otaA | 8048            | 9               | 7611            | 2536|
|                             | otaC    | 1822            | 6               | 1545            | 514|
| (23679 bp)                  | otaB    | 5689            | 2               | 5640            | 1879|
|                             | otaC    | 1822            | 6               | 1542            | 513|
| Aspergillus welwitschiae CBS139.54b | otaA | 8161            | 9               | 7653            | 2550|
|                              | otaB    | 5689            | 2               | 5640            | 1879|
|                              | otaC    | 1833            | 6               | 1545            | 514|
| Aspergillus sulphureus (fresenii) | otaA | 8081            | 9               | 7659            | 2552|
|                              | otaY    | 387             | 1               | 387             | 128|
| (23736 bp)                  | otaB    | 5670            | 2               | 5652            | 1883|
|                              | otaC    | 1827            | 6               | 1545            | 514|
| Aspergillus sclerotiorcarbonarius CBS 121057 | otaA | 8062            | 9               | 7617            | 2538|
|                              | otaY    | 387             | 1               | 387             | 128|
| (19040 bp)                  | otaB    | 5692            | 2               | 5643            | 1880|
|                              | otaC    | 1822            | 6               | 1542            | 513|
| Penicillium nordicum DAOMC 18563 | otaA | 8062            | 9               | 7617            | 2538|
|                              | otaY    | 387             | 1               | 387             | 128|
| (22769 bp)                  | otaB    | 5692            | 2               | 5643            | 1880|
|                              | otaC    | 1822            | 6               | 1542            | 513|
| Penicillium verrucosum BFE808 | otaA | 8113/8065       | 9               | 7662/7614       | 2553/2537|
|                              | otaY    | 387             | 1               | 387             | 128|
| (22931 bp)                  | otaB    | 5689            | 2               | 5640            | 1879|
|                              | otaC    | 1833            | 6               | 1542            | 513|
|                              | otaD    | 1759            | 6               | 1479            | 492|
|                              | otaA    | 8093            | 9               | 7656            | 2551|
|                              | otaY    | 391             | 1               | 391             | 126|
| (23586 bp)                  | otaB    | 5692            | 2               | 5643            | 1880|
|                              | otaC    | 1833            | 6               | 1545            | 514|
| Aspergillus subramanianii CBS 138230 | otaA | 8094            | 9               | 7659            | 2552|
|                              | otaY    | 381             | 1               | 381             | 126|
| 23730 bp                    | otaB    | 5672            | 2               | 5646            | 1881|
|                              | otaC    | 1851            | 6               | 1542            | 513|
|                              | otaD    | 1762            | 6               | 1494            | 497|
|                              | otaA    | 8093            | 9               | 7623            | 2540|
|                              | otaY    | 378             | 1               | 378             | 125|
| (22139 bp)                  | otaB    | 5682            | 2               | 5634            | 1877|
|                              | otaD    | 1746            | 6               | 1494            | 497|
| Aspergillus alliaceus CBS 536.65 | otaA | 8093            | 9               | 7623            | 2540|
|                              | otaY    | 378             | 1               | 378             | 125|
| (22138 bp)                  | otaB    | 5682            | 2               | 5634            | 1877|
|                              | otaC    | 1809            | 6               | 1539            | 512|
|                              | otaD    | 1746            | 6               | 1494            | 497|
|                              | otaA    | 8093            | 9               | 7623            | 2540|
|                              | otaY    | 378             | 1               | 378             | 125|
| (22138 bp)                  | otaB    | 5682            | 2               | 5634            | 1877|
|                              | otaC    | 1809            | 6               | 1539            | 512|
|                              | otaD    | 1746            | 6               | 1494            | 497|

a NRPS, a cytochrome P450 monooxygenase, a basic leucine zipper (bZIP) transcription factor and a halogenase. All of these genes and their associated proteins have been demonstrated to be involved in the biosynthesis of OTA across a range of ochratoxigenic species (Gallo et al., 2012, 2017; Ferrara et al., 2016; Gil-Serna et al., 2018; Wang et al., 2018). Identified and characterized in this study, our genome analyses confirmed the presence of a putative OTA PKS cyclase gene in all currently sequenced OTA producing fungi. The location of the SnoaL cyclase between PKS and NRPS encoding genes as well as its transcriptional direction relative to those genes, was also assessed. Generally, synteny and reciprocal transcription direction of the core cluster genes is conserved, as well as the number of exons for coding sequences of each gene (Figure 1).

Here, we apply the conventional naming system for the genes of OTA cluster that have already been used by Wang et al. (2018). Specifically, it is based on a three-letter code “ota” followed by a capital letter representing each individual gene, that resembles the naming systems usually used for biosynthesis genes of most fungal mycotoxins and secondary metabolites. In the core cluster, the genes encoding for the key proteins PKS and NRPS are named otaA (pks) and otaB (nrps), the genes encoding for the P450 oxidoreductase and the halogenase protein are named otaC (p450) and otaD (hal), and the gene identified in this work and
encoding the putative OTA PKS cyclase is named *otaY* (cyc). For *A. westerdijkiae*, a coding sequence for a predicted PKS cyclase of 128 aa was present between *otaA* and *otaB* genes in the draft annotation of strain CBS112803 at JGI, we therefore included its annotation in the data related to the strain CECT2948 (Gil-Serna et al., 2018). Moreover, the study of OTA cluster in *A. ochraceus* FC1 (Wang et al., 2018) led to the identification of a putative coding sequence for a predicted PKS cyclase of 128 aa, located between *otaA* and *otaB* genes as in the other ochratoxigenic species.
Species of *Penicillium* are also known to produce OTA. In *P. nordicum* strain DAOM 18563, we identified a gene sequence encoding a putative PKS cyclase located between *otaA* and *otaB*. The predicted OTA core cluster gene sequences of *P. verrucosum* strain BFE808 were identified based on homology analysis with a PKS cyclase gene identified between *otaA* and *otaB* as well.

Two other genes encoding a flavin adenine dinucleotide (FAD)-dependent oxidoreductase (*otaE*) and a zinc finger DNA binding protein (*otaR2*) involved in the regulation of some core genes were identified adjacent to the core cluster in *A. ochraceus* and in proximity to the *otaA* gene (Wang et al., 2018). We have found that the genome sequences of a several species encode *otaE* and *otaR2* in the same position. There are exceptions, however: for example, the genomes of *A. sclerotioroniger*, *A. niger* and *A. carbonarius*, have the *otaR2* gene located far from the core cluster. Very few species exhibit only the *otaE* gene in the conserved position close to *otaA*. At the opposite end of the core cluster, several species have genes encoding a peptidase or hydroxylase. The characteristics of all characterized OTA clusters are summarized in Figure 1.

The predicted amino acid sequences of PKSs encoded by *otaA* genes display the characteristic domains of a highly reducing (HR)-PKS (KS, AT, DH, CMet, ER, KR, and ACP) involved in the first step of formation of the pentaketide backbone of the OTA molecule starting from acetate and malonate. Some divergent data has emerged about the length of *otaA* and other OTA genes in *A. niger* strain ATCC13496 used in this study, as compared with data reported by other authors for *A. niger* strain CBS513.88 (Pel et al., 2007; Gil-Serna et al., 2018), the first *A. niger* strain sequenced. Most of these differences are likely due to the variety of automated annotation pipelines used, in particular related to the prediction analysis of exon/intron junctions. Analysis of the strain ATCC13496 in this study led to the identification of a putative PKS cyclase in the *A. niger* species.

In *P. verrucosum* BFE808, two possible start codons were detected for the *otaA* coding sequence, to initiate translation of predicted proteins of 2553 and 2537 aa, respectively. Similarly, two possible start codons were found in the *otaA* open reading frame of *P. nordicum* also. In the case of *A. sclerotioronarius* the analysis of the *otaA* gene revealed the presence of 11 exons instead of 9, as observed in the other species.

The analysis of *otaB*, *otaC*, *otaR1*, and *otaD* genes revealed that they are well conserved in terms of gene length and coding sequence structure in the analyzed species, with the exception of *A. sclerotioronarius* wherein both *otaB* and *otaD* genes exhibited one additional exon compared with other species.

Finally, an unexpected and interesting result came from the analysis of OTA core cluster organization in *A. albertensis* IBT14317 and *A. alliaceus* CBS 536.65. In these two species, the *otaD* gene was found between *otaB* and *otaR1* rather than at the end of the core cluster. It should be noted that while there are slight differences in genome size and gene model content (Kjærboelling et al., 2020) in the revision of Section *Flavi* (Frisvad et al., 2019) *A. albertensis* was tentatively synonymized to *A. alliaceus.*
analyzed resulted related to SnoaL domains of the OTA clusters, while they formed numerous clades, sometimes well supported. On the basis of this first general phylogenetic results evidencing the robustness of OTA cyclases SnoaL domain clustering, we focused our phylogenetic analysis on the 23 cyclases from OTA biosynthetic clusters; a cyclase of *Metarhizium robertsii* found between a PKS and NRPS genes in a partial putative OTA cluster, and the cyclases of *A. parvulus* and *A. cervinus*, closely related to the group of the OTA cyclases. The phylogenetic tree resulting from the Maximum likelihood analysis with the highest log likelihood (−1837.50) is shown in Figure 4. The percentage of trees in which the associated taxa clustered together is shown next to the branches. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. On the same tree the Posterior probability of the Bayesian analysis were also reported on the branch, the Bayesian analysis give the best likelihood tree for run 1 = −1923.89 and for run 2 = −1913.05. The topology of the trees obtained by the two separate analyses is consistent with the Bayesian analysis giving higher support to the OTA cyclase cluster (Figure 4). The result of the phylogenetic analysis clearly confirmed the clustering of the OTA biosynthetic cyclases in a well-supported branch closely related to the *M. robertsii* cyclase and the two cyclase from *A. cervinus* and *A. parvulus*.

**DISCUSSION**

Next-generation sequencing technologies continue to drive a new era in mycology by greatly increasing the catalog of fungal genomes sequences available for analysis. The alacrity with which fungal genomes are being sequenced produces a vast and still-growing amount of information, which must be efficiently integrated and analyzed in order to understand enzymatic and biochemical diversity of these organisms. Despite the development of efficient bioinformatics approaches, maintaining consistency and accuracy in genome curation and annotation remains a daunting task. With each additional OTA producing fungal genome sequence produced, we perform the comparative genomic analysis of OTA biosynthetic cluster. Continuous revision and improvement of the genome annotation, and of comparative analysis of predicted genes allows for refinement and discovery that enhance our knowledge of OTA production and biochemistry. Indeed, our analyses improved the annotation of the OTA cluster in 21 producing species and led to the identification of a previously uncharacterized gene in the cluster. Whole genome analyses have led to the identification of many SM clusters of known metabolites (Kjærbølling et al., 2018), and also to the prediction of novel cryptic clusters for still unknown microbial metabolites (de Oliveira and Ocaña, 2015). But, there are still knowledge gaps associated with the OTA biosynthetic cluster that must be elucidated and new genome data and comparative analysis are critical for accurate annotation and improved biochemical understanding. In this regard, due to its widespread distribution in food, feed and beverages, OTA has been extensively studied in the areas of toxicological, contamination and exposure risks as well as the biochemistry and ecological properties of its producers. Some key enzymatic reactions of OTA biosynthesis have been defined, although the biosynthesis pathway and its regulation have not been completely elucidated. OTA is a hybrid molecule composed of a polyketide dihydroisocoumarin moiety linked via amide bond to the amino acid phenylalanine. A PKS enzyme is involved in the first step.
of formation of the isocoumarin group, starting from acetate and malonate, to originate the characteristic pentaketide skeleton of ochratoxin β (7-carboxy-methyl-mellein), and subsequently a NRPS catalyzes the condensation of the dihydroisocoumarin ring to phenylalanine to form OTB (ochratoxin B). Finally a halogenase enzyme is necessary for the chlorination step that transforms OTB in OTA (Gallo et al., 2012, 2014; Ferrara et al., 2016).

The overall organization of OTA core genes is well conserved among the 21 species included in this study and is in agreement with the evolutionary lineage of ochratoxigenic species. Based on the currently available genome information, our analyses of the core genes of OTA cluster revealed some differences in gene attributes probably due to different bioinformatics pipelines used for genome model prediction. Despite of these discrepancies reported by different authors, the involvement of most of these genes in the biosynthetic pathway of OTA is well established (Gallo et al., 2012, 2014; Ferrara et al., 2016; Wang et al., 2018).

We found that A. steynii, A. muricatus, A. roseglobulosus, A. flocculosus, A. pulvericola, A. westerdijkiae, A. ochraceus, and A. elegans shared the same cluster organization with only minor differences observed in intergenic distance between otaR1 and otaD genes. This group of eight species shared also the same otaR2 gene position, located upstream of the otaA gene adjacent to a FAD-oxidoreductase encoding gene named otaE by Wang et al. (2018). In A. sclerotiniger, A. carbonarius, and A. niger the otaR2 gene is also annotated upstream of the otaA gene, but with more than one predicted gene model located between them. The length of the coding sequence of otaR2 gene is conserved in nearly all the above mentioned species with the exception of A. niger whose coding sequence is considerably shorter than in the others. In the other OTA producer species, the otaR2 gene is not annotated but we believe that this gene has been missed in most of the species due to incomplete annotation of the genomic region adjacent to the core cluster genes. Moreover, upstream of the otaD gene, one to two additional genes are annotated in A. roseglobulosus and A. sulphureus, respectively. Also, in these cases we cannot exclude that their presence needs to be confirmed by further investigations. The analysis of the OTA cluster region of A. albertensis and A. alliaceus reveals a genomic rearrangement wherein a portion of the cluster includes the genes otaC, otaD, and otaR1. The shift of otaD gene near to
OTA gene represents a novelty never observed and described in OTA producing strains. The potential reason for the difference in gene location could be related to several events such as the presence of mobile elements in the proximity of otaD gene. Understanding the differences in OTA gene cluster organization merit further investigation.

Based on the genome annotation of the currently sequenced OTA producer species, we noticed that 17 out of 21 OTA producers presented a putative protein coding for a cyclase (otaY) between the otaA and otaB genes. This gene was not identified in the gene model catalogs of A. welwitschiae, A. elegans, P. nordicum, and P. verrucosum. However, we were able to manually identify and annotate the otaY sequences for inclusion in our phylogenetic analyses. Our phylogenetic analysis supports the hypothesis that the otaY gene is well conserved among the OTA producing species. Interestingly, the preliminary phylogenetic analysis on 863 different SnoaL domains, retrieved from all 225 Aspergillus genomes available in Mycocosm, indicates the clear grouping of the putative OTA cyclase genes within the same phylogenetic group, with a consistent bootstrap. The phylogenetic tree showed that together with the cyclase sequences of the OTA producing strains, a sister clade closely related and well supported, comprises two species not known to produce OTA, namely A. parvulus and A. cervinus. Comparative genomic analysis of these two strains indicated that the OTA cluster was not present, although we noted that the A. cervinus cyclase was neighbored by uncharacterized NRPS and PKS genes. None of the other analyzed SnoaL domains appear to be related to the cyclases putatively involved in OTA biosynthesis, confirming that these latter have likely evolved together following the speciation within Aspergillus genus. The non-OTA associated SnoaL domain containing cyclases formed numerous clades. Thus, a phylogenetic analysis was performed on the restricted group of 23 OTA SnoaL domains (comprising 22 OTA producing species and A. sclerotiorum; not producing OTA but displaying the complete biosynthetic cluster), a cyclase domain of M. robertii in a partial OTA cluster, and the two cyclases domains of A. parvulus and A. cervinus retrieved from the previous analysis. The clustering of 23 OTA SnoaL domains was in accordance with the phylogeny at species and section level for the Aspergillus (Figure 4), with species of section Nigri, section Flavi and Circumdati were well supported. Interestingly, the cyclases of P. verrucosum and P. nordicum clustered together (bootstrap 99/1); but they belong to the well supported clade associated with Aspergillus Sect Circumdati (Figure 4). These findings support the hypothesis of a possible horizontal gene transfer between Aspergillus and Penicillium genera, particularly since it is well known that they come from a common ancestor and belong to the same family of Aspergillaceae (Samson et al., 2014; Houbraken et al., 2020). Moreover, previous data showed high similarity of OTA PKS between Section Circumdati species and P. nordicum (Culebras et al., 2009; Gallo et al., 2014).

In our study, we observed the occurrence of the transcript of the putative cyclase of OTA cluster in three important producing species, confirming that it was expressed during OTA biosynthesis. The characterization of nucleotide and amino acid sequences attested the presence of a domain showing high homology to SnoaLs. SnoaLs proteins have been characterized as belonging to a family of small polyketide cyclases, consisting of approximately 140 amino acids, which catalyze ring closure steps in the biosynthesis of polyketide antibiotics produced in Streptomyces (Sultana et al., 2004). It was suggested that the reaction catalyzed by these enzymes is an intramolecular aldol condensation, by using acid–base chemistry rather than covalent or metal ion catalysis. Despite the structural similarity shared between bacterial and fungal aromatic polyketides, fundamental differences in cyclization strategies exist. It was reported that a PT domain controls the cyclization steps of the nascent non-reducing (NR) fungal polyketides and frequently a thioesterase/Claisen-like cyclase (TE/CLC) is found at the C-terminus of the NR-PKSs to catalyze additional cyclization steps and product release (Zhou et al., 2010). There is limited data regarding cyclization and release processes in the biosynthesis of heterocyclic polyketides produced by fungal highly reducing PKSs (HR-PKSs), like OTA PKS. They are a subgroup of iterative type I fungal PKSs that produce linear, reduced polyketides in an assembly line process. The PKS intermediates remain bound to the megasyme via a thioester linkage during the whole process. Many HR-PKSs lack a C-terminal release domain, even though some HR-PKSs occur as fused dimodules with a C-terminal NRPS module, such as the well-studied lovastatin HR-PKS LovB which contains a C-terminal NRPS condensing (C) domain (Herbst et al., 2018). More often, cyclization is thought to happen spontaneously via intramolecular interactions within the growing polyketide. The alternative hypothesis could be the involvement of a cyclase gene separate from the PKS gene. In this regard, the biosynthetic pathway for aurovertin by the ascomycete Calcarisporium arbuscula is thought to involve a cyclase protein (aurE), with sequence homology to bacterial aromatic polyketide cyclase SnoaL, that may enhance (but is not essential for) pyrone formation and product release from the aurA PKS (Mao et al., 2015; Hang et al., 2016).

Concerning OTA biosynthetic pathway, the cyclization process that leads to the formation of the heterocyclic structure of OTB, in the first stages, has remained unclear. The identification of a gene encoding a SnoaL like protein (otaY) in the OTA cluster suggests its putative role in the cyclization of OTA polyketide backbone.

Our comparative genomic, transcriptomic and phylogenomic analyses support inclusion of the otaY gene in the OTA biosynthetic gene cluster. Moreover, our results pointed out the need for future research to confirm the role of otaY gene in OTA biosynthesis by gene knock-out linked to the analysis of metabolic profile to track the intermediate metabolites accumulated in otaY defective strains compared to wild type strains.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: https://www.ncbi.nlm.nih.gov/genbank/, MT706047; MT706048; and MT706049.
AUTHOR CONTRIBUTIONS

GP, MF, AG, and SB contributed to conception and design of the study. MF and DM performed the experiments and the analysis. AG and MF wrote the first draft of the manuscript and performed the genomic annotation. GP and DM performed the phylogenetic analysis. SB performed the comparative analysis. All authors contributed to the manuscript revision, read and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2020.581309/full#supplementary-material

Supplementary Figure 1 | The highest log likelihood (−118408.77) tree obtained by RAxML analysis of 863 SinoAL domain is shown with bootstrap value. The yellow highlighted cluster represent the 21 SinoAL domains (red cluster) identified in the OTA biosynthetic clusters, plus the sister cluster of two no producing OTA species A. cervinus and A. parvulus.

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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