Twists and Turns in the Salicylate Catabolism of *Aspergillus terreus*, Revealing New Roles of the 3-Hydroxyanthranilate Pathway

Tiago M. Martins, Celso Martins, Paula Guedes, Cristina Silva Pereira

ABSTRACT In fungi, salicylate catabolism was believed to proceed only through the catechol branch of the 3-oxoadipate pathway, as shown, e.g., in *Aspergillus nidulans*. However, the observation of a transient accumulation of gentisate upon the cultivation of *Aspergillus terreus* in salicylate medium questions this concept. To address this, we have run a comparative analysis of the transcriptome of these two species after growth in salicylate using acetate as a control condition. The results revealed the high complexity of the salicylate metabolism in *A. terreus* with the concomitant positive regulation of several pathways for the catabolism of aromatic compounds. This included the unexpected joint action of two pathways—3-hydroxyanthranilate and nicotinate—possibly crucial for the catabolism of aromatics in this fungus. Importantly, the 3-hydroxyanthranilate catabolic pathway in fungi is described here for the first time, whereas new genes participating in the nicotinate metabolism are also proposed. The transcriptome analysis showed also for the two species an intimate relationship between salicylate catabolism and secondary metabolism. This study emphasizes that the central pathways for the catabolism of aromatic hydrocarbons in fungi hold many mysteries yet to be discovered.

IMPORTANCE Aspergilli are versatile cell factories used in industry for the production of organic acids, enzymes, and pharmaceutical drugs. To date, bio-based production of organic acids relies on food substrates. These processes are currently being challenged to switch to renewable nonfood raw materials—a reality that should inspire the use of lignin-derived aromatic monomers. In this context, aspergilli emerge at the forefront of future bio-based approaches due to their industrial relevance and recognized prolific catabolism of aromatic compounds. Notwithstanding considerable advances in the field, there are still important knowledge gaps in the central catabolism of aromatic hydrocarbons in fungi. Here, we disclose a novel central pathway, 3-hydroxyanthranilate, defying previously established ideas on the central metabolism of the aromatic amino acid tryptophan in *Ascomycota*. We also observe that the catabolism of the aromatic salicylate greatly activated the secondary metabolism, furthering the significance of using lignin-derived aromatic hydrocarbons as a distinctive biomass source.

KEYWORDS RNA-seq, aromatic compound catabolism, *Aspergillus nidulans*, *Aspergillus terreus*, 3-oxoadipate pathway, gentisate, tryptophan kynurenine pathway, nicotinate catabolism, secondary metabolism, saprophytic *Ascomycota*
industrial scale (1). The bio-based production of organic acids is currently shifting from food (corn, beet, cane, etc.) to renewable nonfood feedstocks that may include production by engineered microorganisms (2). Aspergilli are used industrially for the production of organic acids (e.g., citric acid and itaconic acid) and secondary metabolites (e.g., the cholesterol-lowering drug lovastatin), hence seen as ideal candidates for novel production processes (3). Lignin, a heterogeneous aromatic plant polymer, is the second most abundant after cellulose. It constitutes a major underutilized residue from various industrial processes (e.g., >100 million tons per year from Kraft pulping alone) (4). Lignin’s high recalcitrance hinders many biotechnological approaches (5), despite the fact that recent progress in biorefinery can ensure its depolymerization into numerous and structurally diverse aromatic hydrocarbons (5, 6). In nature, lignin depolymerization is mostly accomplished by potent oxidative enzymes secreted by fungi that possess multiple strategies for the catabolism of these aromatic hydrocarbons (7). Fungi can use countless peripheral pathways for their catabolism, all of which converge toward a small number of intermediates that undergo ring cleavage in the central pathways. So far, the better described central pathways are of the intermediates catechol, protocatechuate, hydroquinol, homogentisate, and gentisate (8). In the last decade, many of the catabolic genes composing these central pathways have been assigned (8–13). Searching for these genes in the genomes of Dykarinia revealed that the five central pathways are present in both Ascomycota and Basidiomycota, sharing simultaneously high similarity but also potentially some speciation (e.g., a distinctive Basidiomycota protocatechuate 3,4-dioxygenase) (8). In particular, the existence of specific dioxygenases for gallate, hydroquinone, homoprotocatechuate, and pyrogallol, as described for bacteria, remains unclear to date, despite that these compounds are known to be channeled into the known central pathways (10, 14, 15).

Aspergilli generally display in their genomes the orthologous genes for all the better described central catabolic pathways, with the exception of the gentisate pathway, which is either present, partially present, or absent (8, 16). In addition, the gentisate 1,2-dioxygenase orthologous gene is always organized in a cluster that comprises either all the other pathway genes (designated gentisate cluster) or all except the malylpyruvate isomerase gene (designated the “gentisate-like cluster”) (8). The presence of the two clusters in several fungal genomes (e.g., Aspergillus niger) is suggestive of different and unknown functional roles. Aspergillus terreus has a gentisate-like gene cluster in its genome, whereas Aspergillus nidulans does not have a gentisate gene cluster or a gentisate 1,2-dioxygenase orthologous gene. In the present study, we undertook a comparative transcriptome analysis of these two species upon their growth in salicylate as the sole carbon and energy source, using acetate as the control condition. The study was complemented by expression analyses of targeted genes and chemical analysis of the potential intermediates.

RESULTS AND DISCUSSION

Cultivation of Aspergillus terreus in salicylate (20 mM) resulted in the transient accumulation of gentisate up to ca. 3 mM (Fig. 1). This observation conflicts with salicylate catabolism only through the catechol branch of the 3-oxoadipate pathway, as systemically shown in other aspergilli, including in A. nidulans (see reference 9 and references therein). Some yeasts have been shown to use the gentisate pathway for the catabolism of 3-hydroxycinnamates, 3-hydroxybenzoate, and gentisate (2,5-dihydroxybenzoate) (12). The genes of the gentisate pathway are clustered in the genomes of fungi (8), yet in the gentisate-like cluster, the malylpyruvate isomerase gene is absent, as observed in A. terreus (ATEG_06711 to ATEG_06714). This suggests that during the catabolism of salicylate in A. terreus, gentisate catabolism also takes place through a yet unknown pathway. To better understand the catabolism of salicylate in A. terreus, we analyzed its transcriptome response, as well as that of A. nidulans, grown in salicylate. The comparative analysis revealed that both species share a global common response to salicylate compared to the control conditions in acetate (Fig. 2). In the presence of salicylate, 2,703 encoding
genes were found differentially upregulated and 1,489 downregulated in *A. nidulans* and 2,980 were upregulated and 2,328 downregulated in *A. terreus* (see Fig. S1 for principal-component analysis [PCA] and MA plots and Tables S4 and S5 in the supplemental material). The first striking feature is that many of the upregulated genes could not be annotated in KEGG (ca. 80%), while half of the downregulated ones could (ca. 50%) (Fig. 2). This is likely related to the fact that well-known processes/pathways were globally downregulated in salicylate medium, such as carbohydrate metabolism (e.g., tricarboxylic acid [TCA] cycle, glycolysis, and pentose phosphate pathways), energy metabolism (e.g., oxidative phosphorylation) and genetic information processing (e.g., protein processing in the endoplasmic reticulum). As expected, the acetate-utilizing genes (*facA* to *facC*, *acuD* to *acuN*, and *maeA* to *maeB*) were downregulated, including the ones from the glyoxylate shunt (Tables S4 and S5). Globally, the amino acid metabolism was greatly affected in both species. Specifically, the aromatic amino acid metabolism was upregulated and their biosynthesis was downregulated (Tables S4 and S5). The peripheral pathways of the catabolism of aromatic compounds show resemblance to the metabolism of phenylalanine, tyrosine, and tryptophan, suggestive of their upregulation. In agreement, and expectably, the xenobiotics’ biodegradation and metabolism were also upregulated. The analysis of the protein domains demonstrated that those upregulated and enriched are from families of oxidoreductases, hydrolases, transporters, transcriptional regulators, and transferases (Fig. 2). On the contrary, none of the downregulated genes encode abundant enriched protein domain families, with the exception of the flavin adenine dinucleotide (FAD)/NAD(P)-binding domain (IPR023753) found in diverse oxidoreductases (e.g., thioredoxin and glutathione reductases and dihydrolipooyl dehydrogenase).

**Secondary metabolism.** Both *Aspergillus* species grown in salicylate underwent similar transcriptome responses in secondary metabolism (SM), with the upregulation of nearly half of the core biosynthesis genes, whereas only ca. 10% underwent downregulation (Table 1; see Table S3 in the supplemental material). Specifically, in *A. nidulans*, 5 out of 10 laeA-like methyltransferase genes (*llmA*, *llmC*, *llmE*, *llmI*, and *llmJ*) with potential regulatory roles in SM (19) were found to be upregulated, while the remaining five were unaltered. In *A. terreus*, out of the eight laeA-like genes, three were upregulated (ATEG_01440, ATEG_06013, and ATEG_09440), four were unaltered (ATEG_00678, ATEG_01936, ATEG_04864, and ATEG_00912), and only one was downregulated (ATEG_07798). Collectively, the concerted response observed in SM reveals its close relation with the catabolism of the aromatic salicylate.

In salicylate, the monodictyphenone gene cluster of *A. nidulans* was upregulated, whereas that of sterigmatocystin was downregulated, similar to what was reported before using proteome analyses (9). The monodictyphenone biosynthesis relies on
malonyl-coenzyme A (malonyl-CoA) as a precursor (20), formed from plentiful acetyl-CoA made available by the catabolism of salicylate. Besides the monodictyphenone gene cluster, the other known prenyl xanthone biosynthesis genes were also found moderately to highly expressed and upregulated. This included the aromatic DMATS-type prenyltransferase (PTase) genes \textit{xpt}A (AN6784) and \textit{xpt}B (AN12402), the GMC oxidoreductase superfamily gene \textit{xpt}C (AN7998), and the short-chain dehydrogenase/reductase (SDR) gene AN7999. The last was reported before to have no obvious effect on prenyl xanthone biosynthesis (20). Three DMATS-type PTases were also found

FIG 2 Analysis of the RNA-seq data collected for \textit{Aspergillus nidulans} and \textit{A. terreus} grown in salicylate (compared to acetate) comprising functional characterization and protein domains enrichment of the observed differentially expressed genes. KEGG functional categories were slightly simplified by creating super categories that merge the most similar ones. The predicted protein domains of the upregulated and significantly enriched genes in salicylate (\(P \leq 0.05\)) for both species are shown. Fold enrichment and percentage of upregulated genes are displayed in the bottom axis and the log10 \(P\) value in the top axis.

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moderately to highly expressed and upregulated, including those of the nidulanin A cluster (AN11080) and the viridicatin/aspoquinolone cluster (AN11194/AN11202), which use as precursors either kynurenine or anthranilate, and the amino acids phenylalanine and valine, respectively.

In *A. terreus*, salicylate greatly upregulated the asterriquinone and the aspulvinone clusters, comprising the DMATS-type PTases ATEG_09980 and ATEG_01730, respectively. These clusters use as precursors indole-3-pyruvate and 4-hydroxyphenylpyruvate, respectively (21–23), which belong to the tryptophan and the phenylalanine pathways also undergoing upregulation in salicylate. A putative cyclic peptide (ribosomally synthesized and posttranslationally modified peptide [RiPP]) gene cluster underwent massive expression and upregulation in salicylate, of which the gene ATEG_07504 encodes the precursor protein.

Collectively, these findings highlight the interconnection of SM and salicylate catabolism in aspergilli, possibly a consequence of shared regulatory and metabolic routes, including the availability of specific precursors. This deserves further consideration in the near future, especially as the use of simple hydrocarbon aromatics, such as salicylate, can help to disclose unknown biosynthesis clusters and the resultant metabolites.

**Salicylate catabolism in Aspergillus nidulans.** As previously reported, salicylate catabolism in *A. nidulans* most likely results, either directly or via 2,3-dihydroxybenzoate as an intermediate, in the formation of catechol (9), a central substrate for a ring cleavage dioxygenase, which is then channeled to the respective branch of the 3-oxoadipate pathway. This pathway was found mostly upregulated (see Fig. S2 and Table S4 in the supplemental material). However, only the two genes of the non-oxidative decarboxylation pathway and the catechol 1,2-dioxygenase (AN7418, *dhbD*, and AN4532) underwent major expression, while the remaining ones showed only minor expression (AN2114, AN4061, AN4531, and AN10495). We have previously observed that some of these less-expressed genes are not essential for growth in salicylate (9). Accordingly, we previously suggested that the hydroxyquinol variant of the 3-oxoadipate could constitute an alternative path (9). In the present study, we observed by reverse transcription-quantitative PCR (RT-qPCR) expression analyses of the ΔAN4531 mutant grown on salicylate that this alternative path is unlikely (see Fig. S3 in the supplemental material). We also noticed that the dark-colored metabolite (likely 3-oxoadipate enol-lactone that accumulated during the growth of this mutant [9]) disappeared after prolonged growth incubations. These observations suggest that muconate and its catabolic derivatives may be metabolized by broad-substrate-specificity enzymes. The

| Gene regulation | A. nidulans | A. terreus |
|----------------|------------|------------|
| Total No. of SM biosynthesis genes up- or downregulated | 446 | 398 |
| Up | 260 | 118 |
| Down | 39 | 31 |
| Core biosynthesis genes | 96 | 111 |
| PKS | 32 | 32 |
| NRPS | 33 | 39 |
| Hybrid | 1 | 1 |
| PTases | 7 | 11 |
| Terpene | 11 | 10 |
| β-Lactone | 6 | 4 |
| Cyclic peptide | 6 | 13 |
| Siderophore | 0 | 1 |

*PKS, polyketide synthase; NRPS, nonribosomal peptide synthetase; Hybrid, PKS and NRPS; and PTases, aromatic prenyltransferases.

*Genes were considered upregulated (Up) when the fold change was >1, downregulated (Down) with a fold change of <−1, and both when the P value was <0.05.

**Table 1** Observed regulation for secondary metabolism biosynthesis genes in salicylate medium

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Table 1 summarizes the observed regulation of secondary metabolism biosynthesis genes in salicylate medium. The table lists the number of genes up- or downregulated in *A. nidulans* and *A. terreus* for each of the main biosynthesis clusters and core genes. *A. nidulans* showed a total of 446 genes upregulated and 39 downregulated, while *A. terreus* had 398 upregulated and 51 downregulated genes. The table highlights the differences in gene expression between the two species, with *A. nidulans* showing a higher number of genes overall.

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upregulation of several oxidoreductase and hydrolase gene families (Fig. 2) most likely contributes to redundancy in the metabolism of muconate and may offer metabolic plasticity in similar contexts. Besides the two salicylate monooxygenase genes of the catabolic pathway (AN2114 and AN7418), 45 other aromatic-ring FAD-binding monooxygenase genes (IPR002938) were found moderately to highly expressed and upregulated, including seven phenol monooxygenase genes (IPR038220). A major facilitator superfamily (MFS) transporter gene (AN10845) that underwent upregulation in salicylate is located contiguous to dhbD. These two genes show shared synteny within aspergilli and penicillia, suggestive that this MFS transporter can be a 2,3-dihydroxybenzoate transporter. This requires validation since numerous other genes of this superfamily were also positively regulated.

One remarkable observation is that we found also a cluster of four genes highly expressed and upregulated in salicylate. It comprises the following genes: the putative salicylate 1-monooxygenase (AN2114) previously identified by us (9), a domain of unknown function (DUF3425) gene (AN2115), a NAD-dependent epimerase/dehydratase (AN2116), and an amidohydrolase (AN2118) (Table S4). The last gene has sequence homology to orsB (orsellinic cluster) and dhbD, both encoding enzymes that catalyze the decarboxylation of aromatic hydrocarbons. This cluster is only present in the genomes of aspergilli belonging to the Nidulantes section (see Fig. S4 in the supplemental material), being absent in most aspergilli, including A. terreus, hence suggestive of a high specific functional role. One hypothesis is that this putative salicylate 1-monooxygenase plays other functions besides the hydroxylation of salicylate in this fungus, possibly acting in other peripheral pathways or in secondary metabolism.

**Salicylate catabolism in Aspergillus terreus.** The known pathways in microorganisms for the catabolism of salicylate involve either the catechol branch of the 3-oxoadipate pathway or the gentisate pathway (7). We observed in A. terreus that salicylate was hydroxylated to 2,3-dihydroxybenzoate, similarly to what occurs in A. nidulans, but also to gentisate in significant amounts (Fig. 1). The orthologous or predicted genes for these pathways are present in the genome of A. terreus. The transcriptome sequencing (RNA-seq) data revealed that these genes were mostly unaltered in salicylate medium at the respective time point (Fig. 3; Table S5). However, massive upregulation of the non-oxidative decarboxylase gene (ATEG_06350) and key genes coding in the catechol branch (ATEG_09602 and ATEG_03095) took place at early time points during the incubation in salicylate medium, as verified by their RT-qPCR expression levels (Fig. 4). Corroborating this result, muconolactone, cis,trans-muconate, and 2,3-dihydroxybenzoate (Fig. 1) were detected in small amounts in the extracellular medium. Small amounts of hydroquinone and methyl ester conjugates of salicylate, gentisate, and hydroquinone were detected as well (see Fig. S5 in the supplemental material). An additional compound was noticed showing characteristics that suggest it constitutes an anthranilate derivative, regardless of the fact that its precise identity remains doubtful (Fig. S5).

The upregulation of the gentisate-like cluster (ATEG_06711 to ATEG_06714) was less expressive and occurred at later time points compared to the catechol branch (Fig. 4). The salicylate 3-monooxygenase (AN7418) orthologous gene is not present in the genomes of many aspergilli, including A. terreus. In addition, the identity of salicylate 5-monooxygenase is even more obscure as it has not been previously reported in fungi. As observed here for A. nidulans, 32 aromatic-ring FAD-binding monooxygenases (IPR002938) were found moderately to highly expressed and upregulated, including six phenol monooxygenases (IPR038220), hindering a precise assignment of the salicylate monooxygenases at this stage.

The RNA-seq data showed the upregulation of several genes of the phenylalanine, tyrosine, and tryptophan catabolism, as well as three gene clusters, namely, the homogentisate cluster (ATEG_05922 to ATEG_05926), the nicotine-inducible cluster (ATEG_08614 to ATEG_08620 and ATEG_05673), and a cluster unknown in fungi (Fig. 3; Table S5). The first two have been previously described in A. nidulans (24, 25). The last
cluster is here described for the first time in fungi, which we designate as the 3-hydroxyanthranilate catabolic cluster. Three of the comprised genes of this cluster putatively encode three consecutive enzymes of the tryptophan kynurenine pathway (26), namely, 3-hydroxyanthranilate dioxygenase (3HAO; ATEG_10009-10; not annotated in FIG 3 Diagram of the metabolism of salicylate in Aspergillus terreus. The differentially expressed genes’ (RNA-seq) regulation and abundance are represented by circles of various sizes and colors (see the inset legend). Triangles and squares indicate identification by liquid chromatography (LC) and GC-TOF MS, respectively.
the public databases [see Table S2 in the supplemental material]), aminomuconate semialdehyde dehydrogenase (AMSDH; ATEG_10010), and 2-amino 3-carboxymuconate 6-semialdehyde decarboxylase (ACMSD; ATEG_10009). This cluster, as well as some upstream pathway genes, was observed to be upregulated in salicylate (Fig. 3; Table S5). Three types of indoleamine 2,3-dioxygenases (IDO) present in aspergilli can catalyze L-tryptophan to kynurenine at the start of the pathway, although with different biochemical characteristics and unknown functional roles (27). IDOα (ATEG_03482) was found downregulated and highly expressed, while IDOβ (ATEG_00759) and IDOγ (ATEG_01829) were found upregulated and with even higher expression levels. However, a fourth IDO gene (ATEG_07359) is present in the genome of A. terreus, which exhibited almost no expression in salicylate and acetate media. The 3-hydroxyanthranilate dioxygenase genes in fungi have been merely associated with an alternative path for NAD⁺ biosynthesis (28). On the contrary, our data support the existence of the catabolic tryptophan kynurenine pathway forming 2-aminomuconate in fungi (Fig. 3 and 4),

FIG 4 Expression levels of genes coding in the metabolic pathways of Aspergillus terreus in salicylate medium along cultivation (gray bars represent 55, 75, and 90 h and the last time point used for RNA-seq) compared to the control in acetate medium (white bars represent the time point used for RNA-seq). The central intermediates of each metabolic pathway are indicated. Values represent means ± standard errors of the means from three biological replicates: *, P < 0.05; and **, P < 0.005.
regardless of the fact that none of the intermediates of this pathway (e.g., 3-hydroxyanthranilate) could be definitely identified (Fig. S5). Under the cultivation conditions used here, also catechol did not accumulate, hindering its detection, similar to what was reported before (9). The reaction product of 3-hydroxyanthranilate dioxygenase is 2-amino-3-carboxymuconate semialdehyde, which can be converted enzymatically to 2-aminomuconate semialdehyde or decay nonenzymatically to quinolinate, which is the precursor of NAD$^+$ (Fig. 3). Finally, 2-aminomuconate semialdehyde can also decay non-enzymatically to picolinate, a bidentate chelating agent, of which the metabolism remains largely unknown (26).

The 3-hydroxyanthranilate catabolic gene cluster is present across Ascomycota, including some species of Aspergillus, Fusarium, Metarhizium, and Penicillium (Fig. 5), but absent in Saccharomycotina. In some aspergilli, two similar clusters are present (e.g., in Aspergillus oryzae), with the second cluster showing structural conservation and higher sequence homology to the cluster found in Metarhizium and Penicillium;
indicative of a more recent horizontal gene transfer event. The cluster is absent in A. niger and Fusarium graminearum, where L-tryptophan catabolism proceeds via anthranilate, 2,3-dihydroxybenzoate, and the catechol branch of the 3-oxoadipate pathway (29, 30). A fourth gene of the YigF/YER057c/UK114 family (IPR006175) is recurrently present in the cluster, but not in A. terreus (Fig. 5). Proteins of this diverse family are expected to have various metabolic roles extending from endoribonuclease activity and 2-aminomuconate deaminase to enamine/imine deaminase of the branched-chain amino acid biosynthesis pathway (31). Its orthologous gene in A. terreus (ATEG_05745) was found moderately expressed and upregulated in salicylate (Fig. 3 and 4): hence, possibly it encodes a 2-aminomuconate deaminase. It has been shown that this deamination reaction may also occur nonenzymatically, forming the same products: 2-hydroxymuconate and its tautomer 2-oxo-3-hexenedioate (32). In line with this, herein we noticed that a putative 4-oxalocrotonate tautomerase gene (ATEG_05150) was found upregulated, suggestive that the encoded protein (IPR014347) may mediate 2-oxo-3-hexenedioate formation (Fig. 3). This compound may undergo further reduction to 2-oxoadipate or decarboxylation and hydration to 4-hydroxy-2-oxopentanoate. The subsequent decarboxylation of 2-oxoadipate to glutaryl-CoA by the oxoglutarate dehydrogenase complex (encoded by ATEG_03911 and ATEG_09000) could not be verified by the RNA-seq data. We observed that the TCA cycle in A. terreus (and in A. nidulans) was found highly downregulated in salicylate medium, and the predicted glutaryl-CoA dehydrogenase (ATEG_08866) was also found downregulated (Table S5).

Several genes of the leucine catabolic pathway were found extremely expressed and upregulated in salicylate, namely, the E1 and E2 components of the branched-chain α-keto acid dehydrogenase complex (ATEG_03862, ATEG_06635, and ATEG_09087), isovaleryl-CoA dehydrogenase (ATEG_06574), and 3-methylcrotonyl-CoA carboxylase (ATEG_06573 and ATEG_06576). However, the remaining genes of the leucine catabolism were downregulated or only moderately expressed (Table S5). Taken together, it remains unclear if those genes that underwent upregulation have a role in the catabolism of the branched-chain 2-oxoacid 4-hydroxy-2-oxopentanoate, namely, in its decarboxylation to 3-hydroxybutyryl-CoA. The predicted 3-hydroxybutyryl-CoA dehydrogenase (ATEG_02736) and the downstream acetyl-CoA acetyltransferase (ATEG_00012) were found downregulated, although highly expressed (Fig. 3). Importantly, the catabolism of branched-chain amino acids was previously shown to be upregulated under conditions where 2-hydroxymuconate was produced (33).

In aspergilli, the 3-hydroxyanthranilate catabolic gene cluster shows synteny to genes likely coding in the peripheral pathways (Fig. 5), some of which upregulated by salicylate (ATEG_10004, ATEG_10005, and ATEG_10008). Two of these genes, encoding an aryl-CoA ligase (AMP forming) (ATEG_10005) and an SDR (ATEG_10008), show low homology to characterized proteins. However, the FAD-binding monoxygenase ATEG_10004 gene, which here underwent major expression, contains the sequence motif of a Baeyer-Villiger monoxygenase (34), a distinct class of flavoproteins that catalyze the insertion of an oxygen atom in a C–C bond of a ketone forming an ester. In addition, another FAD-binding monoxygenase gene (ATEG_10014) that was found greatly expressed and upregulated shows discrete homology to the salicylate 1-monoxygenase (decarboxylating) gene, which is located in the vicinity of the 3-hydroxyanthranilate catabolic cluster (Fig. 5).

The six orthologous genes of the A. nidulans nicotinate-inducible gene cluster (25) are part of a highly expressed and upregulated gene cluster of nine genes in A. terreus grown in salicylate that includes three genes not yet described: two FAD-dependent monoxygenase genes (ATEG_08614 and ATEG_08615b) and an SDR gene (ATEG_08615a) (Fig. 6). ATEG_08614 shows discrete homology to the bacterial 6-hydroxynicotinate 3-monoxygenase (Q8BFY2) that forms 2,5-dihydroxypyridine as part of nicotinate catabolism, but also to 3-hydroxybenzoate 6-monoxygenase (Q9F131) and ornellinic acid 1-monoxygenase (J4VWM7). ATEG_08615b is a phenol hydroxylase (IPR012941). Their major coregulation in salicylate is suggestive of participation in the nicotinate metabolism.
In addition, its cluster preservation in aspergilli and across Ascomycota, being however divided into two clusters in the genome of *A. nidulans* (Fig. 6), strengthens the hypothesis of their role in the nicotinate metabolism. Expanding the cluster analysis across numerous fungal species (Fig. 6) revealed the presence of two additional colocalized genes—an *ad e a c e-tylase gene* (*ATEG_05673*) and an *amidase gene* (*ATEG_05674*)—that are not located in the nicotinate-inducible cluster in the majority of aspergilli, including in *A. terreus* and *A. nidulans*.
nidulans. In salicylate, only the deacetylase gene (ATEG_05673) was found highly expressed and upregulated, consistent with its assignment to the nicotinate metabolism. In general, the nicotinate-inducible genes of A. nidulans showed to be somehow upregulated in salicylate medium, although to a much lesser extent compared to A. terreus (Fig. 6). This together with the moderate co-upregulation of the single-copy 3-hydroxyanthranilate dioxygenase (3HAO) gene (AN11252, orthologous to ATEG_00088) in A. nidulans (Table S4) pushes forward the idea of an alternative pathway for the metabolism of aromatic compounds. The existence of such a metabolic route possibly explains why the metabolism of the quorum-sensing molecule phenylethyl alcohol yielded the pyridine ring of NADH and NADPH in Candida albicans (35).

The gentisate-like cluster contains a FAD-binding monooxygenase gene (ATEG_06711) that did not undergo initial upregulation (Fig. 4). A similar trend was noticed for the gentisate 1,2-dioxygenase orthologous gene (ATEG_06712). These observations exclude their foreseen involvement in the hydroxylation of salicylate to gentisate and its subsequent ring cleavage. We have identified another gentisate 1,2-dioxygenase gene (ATEG_07116-17), which is not currently annotated (Table S2), that showed no altered expression levels during cultivation in salicylate (Fig. 4). Both A. terreus and A. nidulans were unable to grow at the expense of gentisate (5 or 10 mM), regardless of the capacity to cometabolize gentisate (depletion of 5 mM in less than 3 days with acetate-pregrown cultures). In addition, gentisate may also be transformed to hydroxyquinol by salicylate 1-monooxygenase (36). However, the expression analysis of the two hydroxyquinol 1,2-dioxygenase genes (ATEG_00629 and ATEG_07465), orthologous of the two recently biochemically characterized genes of A. niger (11), as well as of the predicted maleylacetate reductase gene (ATEG_03722), showed no great alteration (Fig. 4; Table S5). The mild upregulation of the hydroxyquinol pathway is more likely to be related to the detection of hydroquinone (Fig. S5), the latter resulting from non-oxidative decarboxylation of gentisate (Fig. 3).

The homogentisate pathway for the catabolism of phenylalanine and tyrosine (chorismate as a common precursor) was found mostly upregulated in salicylate (Fig. 3 and 4). Several routes can be used for the metabolism of their corresponding precursors, phenylpyruvate and 4-hydroxyphenylpyruvate, likely involving a small group of enzymes capable of mediating analogous reactions (37). In line with this idea, we observed that a pyruvate decarboxylase PDC1-like gene (ATEG_06881) and several alcohol and aldehyde dehydrogenase genes (Table S5), potentially coding for enzymes mediating the production of fusel alcohols and acids (e.g., tryptophan to tryptophol) (38), underwent upregulation in salicylate. These observations suggest that the Ehrlich pathway was likely upregulated, further reinforced by the concurrent upregulation of phenylacetate 2-hydroxylase and 3-hydroxyphenylacetate 6-hydroxylase orthologous genes (ATEG_09785 and ATEG_08499) encoding monooxygenases of the fusel acids phenylacetate and 4-hydroxyphenylacetate or their products.

**Conclusions.** Analysis of the transcriptome associated with the catabolism of salicylate in two species of Aspergillus showed its intimate relationship with the secondary metabolism, which was particularly obvious for metabolites requiring action of aromatic prenyltransferases. Exogenous aromatic compounds and their metabolites can provide the building blocks for the synthesis of many secondary metabolites, as well established, e.g., for penicillin (i.e., phenylacetate) (37). We observed that the metabolism of salicylate yields gentisate in A. terreus, likely also producing gentisyl alcohol, an intermediate of secondary metabolites such as patulin (39, 40). This observation suggests that usage of salicylate (or gentisate) as a carbon source may improve the production of target metabolites. Salicylate also upregulated several other catabolic pathways in A. terreus, including the tyrosine metabolism that may provide plentiful SM’ precursors. Tyrosine catabolic genes and secondary metabolite core biosynthetic genes showed similar regulation in penicillia (41). Understanding the relationship between salicylate and SM in fungi may provide unexpected findings on plant-fungus...
interaction, specifically how fungi counteract salicylate-dependent plant defense mechanisms (42).

Collectively our data showed the high complexity of the metabolism of salicylate in A. terreus with concomitant upregulation of several pathways for the catabolism of aromatic compounds at distinct timings (Fig. 4). On the contrary, A. nidulans used only a major route (Fig. S2), similar to that previously reported for many fungal species (see reference 9 and references therein). One unexpected idiosyncrasy of A. terreus is the formation of significant amounts of gentisate in the salicylate medium, and its subsequent metabolism through pathways not yet fully disclosed. However, it is clear that gentisate metabolism led to upregulation of the homogentisate pathway, as well as that of the nicotinate catabolism and 3-hydroxyanthranilate catabolic pathways, none of which could be anticipated. This idea is strengthened because in A. nidulans, which did not yield gentisate through salicylate catabolism, these pathways remained mostly unregulated (or some are even absent). The joint action of the nicotinate metabolism and the 3-hydroxyanthranilate catabolic pathway in A. terreus should be seen as a novel route for the degradation of aromatic compounds. One of the most striking observations is that the gentisate-like gene cluster in this fungus is apparently not involved in the catabolism of the formed gentisate, suggestive of a yet unknown role. In addition, 2-hydroxymuconate is one possible intermediate of the 3-hydroxyanthranilate catabolic pathway, as observed elsewhere in pyrogallol catabolism (43), possibly denoting a convergence of the two pathways. Naturally, this may provide a straightforward strategy for solving the assignment of the genes associated with the catabolism of 2-hydroxymuconate in fungi.

To date, the catabolism of tryptophan in fungi was thought to involve anthranilate and the catechol branch of the 3-oxoadipate pathway. On the contrary, our data on A. terreus and the presence of the 3-hydroxyanthranilate catabolic pathway genes in some Ascomycota genomes suggest that the tryptophan kynurenine pathway in these species shares similarity with that of higher eukaryotes. This opens the exciting possibility of using fungi as model organisms for the investigation of diseases related to tryptophan catabolism: e.g., study of the toxic effects of quinolinate in the progress of Huntington’s disease (26). The 3-hydroxyanthranilate catabolic pathway shows lower occurrence in Ascomycota genomes than the other central aromatic catabolic pathways. Why a smaller number of Ascomycota possess this catabolic plasticity raises intriguing questions about its role and evolutionary traits, especially as some aspergilli have two similar 3-hydroxyanthranilate catabolic gene clusters.

Finally, the sustainable bio-based production of organic acids requires the usage of renewable nonfood biomass, with lignin moving toward the pole position. The transformation of lignin-related aromatic monomers into value-added chemicals may be conceived by the so-called “biological funneling” (5). To date, a delocalized cytosol biosynthesis of TCA cycle intermediates (3) has been used for their production from sugars. Since the aromatic hydrocarbon catabolism in fungi occurs also at the cytosol (44, 45), it is possible to foresee engineered fungal strains as tools to convert these compounds into central intermediates of the catabolism of aromatics and subsequently into organic acids. Our study provides another piece of the puzzle of the catabolism of aromatics in fungi, revealing new catabolic paths (some unexpected) and their putative genes, and proposes future paths to further dissect their functional roles, evolutionary traits, and technological relevance.

MATERIALS AND METHODS

**Strains and growth conditions.** Aspergillus nidulans FGSC A4 or A1147 and A. terreus FGSC A1156 asexual spores were harvested and maintained as frozen suspensions at -80°C (46). Cultures were initiated with 10⁶ spores/ml and incubated with orbital agitation (250 rpm) in the dark at 37°C. Batch cultures were performed in 250-ml Erlenmeyer flasks with a working volume of 50 ml. A low-nitrogen minimal medium was used containing per liter 3 g NaNO₃, 0.01 g ZnSO₄·7H₂O, 0.005 g CuSO₄·5H₂O, 0.5 g MgSO₄·7H₂O, 0.01 g FeSO₄·7H₂O, and 0.5 g KCl. Filter-sterilized salts were added to an autoclave-sterilized 100 mM potassium phosphate (pH 6.0) solution. The carbon sources were added either directly to the phosphate solution for sterilization (60 mM sodium acetate [control]) or to the mineral medium after
filter sterilization (20 mM sodium salicylate). Solid minimal medium was jellified with 5 g/liter agarose, and cultures were grown at 30°C in the dark, on top of a polyvinylidene difluoride (PVDF) membrane (Hybond-P, 0.45-μm pore; GE Healthcare) (47).

Metabolite identification and quantification. Filtrated culture media and the ensuing ethyl acetate extracts (1:1 [vol/vol]; ca. 50-fold concentrated) were chromatographically analyzed for the identification and/or quantification of the aromatic compounds and their putative intermediates either by ultra-performance liquid chromatography (UPLC) (48) or by high-performance liquid chromatography (HPLC). HPLC was performed using a Symmetry C_{18} column, 5 μm, 4.6 mm by 250 mm (Waters Corporation, USA), set at 26°C. The mobile phase, at a flow rate of 0.9 ml min⁻¹, consisted of 0.1% trifluoroacetic acid (solvent A) and acetonitrile (solvent B), set as follows: a linear gradient of 0.5 to 100% B in 30 min, followed by 10 min at 100% B, with 2 min to return to the initial conditions and 10 min to reequilibrate the column. To guide the chromatographic identifications, the standards were used: salicylic acid, 2,3-dihydroxybenzoic acid, gentisic acid, catechol, anthranilic acid, 3-hydroxyanthranilic acid, and cis, trans-muconic acid. The ethyl acetate extracts were also analyzed by gas chromatography-time of flight mass spectrometry (GC-TOF MS) with a Pegasus BT mass spectrometer equipped with an L-PAL3 automatic sampler (LECO). The system was operated using the ChromaTOF software (v5.40.12.0.60635) for Pegasus BT. The separation of the analytes was carried out using a DB-SMS Ultra Inert capillary column (25 m by 0.250 mm by 0.24 μm; Agilent). The carrier gas used was helium at a flow rate of 1.0 ml/min. The split ratio for the injector was set to 1:ÅÅ, with a total injection volume of 1 μl. The front inlet and ion source were both kept at 250°C and the transfer line at 290°C. Oven temperature was set to equilibrate at 60°C for 1 min, before initiation of sample injection and GC run. Following sample injection, oven temperature was maintained at 60°C for another 1 min, increased at a rate of 5°C/min to 290°C, and held for 5 min. The mass detection was operated in electron ionization (EI) mode (70 eV) with a detector voltage of 1,800 V. Full scan mode with a mass range of m/z 40 to 600 and acquisition rate of 15 Hz was used as data acquisition method. Acquisition delay was set at 420 s. Chromatogram data acquisition, baseline correction, peak deconvolution, analyte alignment, peak area integration, and analyte identification by mass spectral searches (based on National Institute of Standards and Technology and Wiley libraries) were performed using the LECO ChromaTOF. Peaks with a similarity of 80% or more and a maximum relative Kovats index deviation of 10% were assigned putative by-product identities based on the mass spectral libraries. The analyses focused on those peaks present in all the salicylate-derived extracts but absent in the control (acetate medium ethyl acetate extracts), relying on the spectral features of pure standards whenever possible.

Transcriptome analysis. Total RNA extraction of liquid-grown mycelia was performed essentially as previously described (46), using a TissueLyser LT (Qiagen) for cell disruption. The quantity and quality of RNA were determined by capillary electrophoresis using the HS RNA kit and 5200 Fragment analyzer (Agilent Technologies). For single-end RNA sequencing (RNA-seq), libraries were generated using the Smart-Seq2 mRNA assay (Illumina, Inc.) according to the manufacturer’s instructions. Twelve samples were indexed and sequenced on the Illumina NextSeq550 (20 million reads per sample). Generated FastQ files were analyzed with FastQC, and any low-quality reads were trimmed with Trimmomatic (49).

All libraries were aligned to the corresponding model fungus, either A. nidulans FGSC A4 genome assembly (ASM149202v2) or A. terreus NIH2624 genome assembly (ASM14961v1) with gene annotations from Ensembl Fungi v.37 using HISAT2 v.2.1.0 (50) and only matches with the best score were reported for each read. All RNA-seq experiments were carried out in three biological replicates. Differential expression analysis was performed using DESeq2 v.1.24.0 (51). The principal-component analysis (PCA) plot and MA plots were also generated using the DESeq2 package. The genes that showed more than log 1-fold expression changes with an adjusted P value of <0.05 are defined as significantly differentially expressed genes in this analysis. Transcript abundance was defined as the number of reads per kilobase of transcript per million mapped reads (RPKM). The full genomes were scanned using InterProScan v.5.32-71.0 (52) to build a protein domain database for further enrichment analyses. The enrichment analyses were performed with the FunRich software v.3.1.3 (53), using the constructed protein domain databases as the terms library and the hypergeometric test with a P value of <0.05.

Reverse transcription-quantitative PCR analysis. Total RNA extraction was performed as reported above, and cDNA synthesis was performed as previously described (46). Oligonucleotide pairs were designed using Primer-BLAST (54) and supplied by STAB Vida (Oeiras, Portugal) (see Table S1 in the supplemental material). The reverse transcription-quantitative PCR (RT-qPCR) analysis was performed in a CFX96 thermal cycler (Bio-Rad), using the SsoFast EvaGreen Supermix (Bio-Rad), 250 mM each oligonucleotide, and the cDNA template equivalent to 10 ng of total RNA, at a final volume of 10 μl per well, in three biological replicates. The PCR conditions were enzyme activation at 95°C for 30 s; 40 cycles of denaturation at 95°C for 5 s and annealing/extension at 60°C for 15 s, and melting curve obtained from 65 to 95°C, consisting of 0.5°C increments for 5 s. Data analyses were performed using the CFX Manager software v.3.1 (Bio-Rad). The expression of each gene was normalized by the expression of the 60S ribosomal protein L33-A gene, AN2980 or ATEG_01624. The expression of all target genes was normalized by the expression of the 60S ribosomal protein L33-A gene, AN2980 or ATEG_01624. The expression of all target genes was normalized as the number of reads per kilobase of transcript per million mapped reads (RPKM). The full genomes were scanned using InterProScan v.5.32-71.0 (52) to build a protein domain database for further enrichment analyses. The enrichment analyses were performed with the FunRich software v.3.1.3 (53), using the constructed protein domain databases as the terms library and the hypergeometric test with a P value of <0.05.

Gene structure prediction. Automated gene structure prediction of new genes or of poorly annotated genes was done using FGENSEW with or without similar protein-based gene prediction (55). Gene structure was manually curated (Table S2) and used in the transcriptome analysis (52).

Identification of gene clusters. Metabolic gene clusters were identified within annotated genomes through sequence homology searches using MultiGeneBlast v.1.11.13 (56). A recent customized database created from online GenBank entries was done using the same software. Identification of secondary metabolite biosynthesis gene clusters was done using antiSMASH v.5.0 (57), and the results were
incorporated into available information (see Table S3 in the supplemental material), namely, from pre-computed results at MycoCosm obtained by using SMURF [58, 59].

**Data availability.** Transcript RPKM values of differentially expressed genes are contained in Tables S4 and S5, and raw data with metadata were deposited in the Sequence Read Accession no. PRJNA612036.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**FIG S1**, PDF file, 0.3 MB.

**FIG S2**, TIF file, 0.5 MB.

**FIG S3**, TIF file, 0.7 MB.

**FIG S4**, TIF file, 0.3 MB.

**FIG S5**, TIF file, 1.1 MB.

**TABLE S1**, PDF file, 0.5 MB.

**TABLE S2**, PDF file, 0.5 MB.

**TABLE S3**, PDF file, 0.5 MB.

**TABLE S4**, XLSX file, 0.8 MB.

**TABLE S5**, XLSX file, 1 MB.

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**REFERENCES**

1. Becker J, Lange A, Fabarius J, Wittmann C. 2015. Top value platform chemicals: bio-based production of organic acids. Curr Opin Biotechnol 36:168–175. https://doi.org/10.1016/j.cojbio.2015.08.022.

2. Skoog E, Shin JH, Saez-Jimenez V, Mapelli V, Olsson L. 2018. Biobased adipic acid—The challenge of developing the production host. Biotechnol Adv 36:2248–2263. https://doi.org/10.1016/j.biotechadv.2018.01.012.

3. Karaffa L, Kubicek CP. 2019. Citric acid and itaconic acid accumulation: variations of the same story? Appl Microbiol Biotechnol 103:2889–2902. https://doi.org/10.1007/s00253-018-9060-7.

4. Becker J, Wittmann C. 2019. A field of dreams: lignin valorization into chemicals, materials, fuels, and health-care products. Biotechnol Adv 37:107360.https://doi.org/10.1016/j.biotechadv.2019.02.016.

5. Rodriguez A, Salvachúa D, Katahira R, Black BA, Cleveland NS, Reed M, Smith H, Baidoo EEK, Keasling JD, Simmons BA, Beckham GT, Gladden JM. 2017. Base-catalyzed depolymerization of solid lignin-rich streams enables microbial conversion. ACS Sustainable Chem Eng 5:8171–8180. https://doi.org/10.1021/acssuschemeng.7b01818.

6. Shuai L, Amiri MT, Questell-Santiago YM, Héroguel F, Li Y, Kim H, Meilan R, Chapple C, Raphael J, Luterbacher JS. 2016. Formaldehyde stabilization facilitates lignin monomer production during biomass depolymerization. Science 354:329–333. https://doi.org/10.1126/science.aaf7810.

7. Lubbers RJM, Dilokpimol A, Visser J, Makelä MR, Hilden KS, de Vries RP. 2019. A comparison between the homocyclic aromatic metabolic pathways from plant-derived compounds by bacteria and fungi. Biotechnol Adv 37:107396. https://doi.org/10.1016/j.biotechadv.2019.05.002.

8. Martins TM, Martins C, Silva Pereira C. 2019. Multiple degrees of separation in the central pathways of the catabolism of aromatic compounds in fungi belonging to the Dikarya sub-kingdom. Adv Microb Physiol 75:177–203. https://doi.org/10.1017/amus.2019.07.003.

9. Martins TM, Hartmann DO, Planchon S, Martins I, Renaut J, Silva Pereira C. 2015. The old 3-oxoadipate pathway revisited: new insights in the catabolism of aromatics in the saprophytic fungus Aspergillus nidulans. Fungal Genet Biol 74:32–44. https://doi.org/10.1016/j.fgb.2014.11.002.

10. Gerecsová G, Neboháčková M, Zeman I, Pyszcz L, Tomáška L, Gabaldon T, Nosek J. 2015. Metabolic gene clusters encoding the enzymes of two branches of the 3-oxoadipate pathway in the pathogenic yeast Candida albicans, FEMS Yeast Res 15:fov006. https://doi.org/10.1093/femsyr/fov006.

11. Semana P, Powolowsi J. 2019. Four aromatic intradiol ring cleavage dioxygenases from Aspergillus niger. Appl Environ Microbiol 85:e01786-19. https://doi.org/10.1128/AEM.01786-19.

12. Holesová Z, Jakubková M, Zavadilakova I, Zeman I, Nosek J. 2011. Gentisate and 3-oxoacid pathways in the yeast Candida parapsilosis: identification and functional analysis of the genes coding for 3-hydroxybenzoate 6-hydroxylase and 4-hydroxybenzoate 1-hydroxylase. Microbiology (Reading) 157:2152–2163. https://doi.org/10.1099/micro.0.048213-0.

13. Mingot JM, Penalva MA, Fernandez-Canon JM. 1999. Disruption of phacB, an Aspergillus nidulans gene encoding a novel cytochrome P450 monooxygenase catalyzing phenylacetate 2-hydroxylation, results in penicillin overproduction. J Biol Chem 274:14545–14550. https://doi.org/10.1074/jbc.274.21.14545.

14. Patel TR, Hameed N, Armstrong S. 1992. Metabolism of gallate in the pathogenic yeast Aspergillus niger. Yeast 8:440–446. https://doi.org/10.1002/yea.3250080403.

15. Ferrer-Sevillano F, Fernandez-Canon JM. 2007. Novel phacB-encoded cytochrome P450 monooxygenase from Aspergillus nidulans with 3-hydroxyphenylacetate 6-hydroxylase and 3,4-dihydroxyphenylacetate 6-hydroxylase activities. Eur J Lipid Sci Technol 109:514–520. https://doi.org/10.1002/ejlt.200700226-06.

16. Greene GH, McGary KL, Rokas A, Slot JC. 2014. Ecology drives the
35. Han T-L, Theisen JM, Duran RM, Grayburn WS, Calvo AM, Keller NP. 2013. Secondary metabolism and development is mediated by LmFC control of VeA subcellular localization in "Aspergillus nidulans." PLoS Genet 9: e1003193. https://doi.org/10.1371/journal.pgen.1003193.

36. Sanchez JP, Entwistle R, Hung JH, Yae-gash J, Jain S, Chiang YM, Wang CC, Oakley BR. 2011. Genome-based deletion analysis reveals the prenyl xanthone biosynthesis pathway in "Aspergillus nidulans." J Am Chem Soc 133:4010–4017. https://doi.org/10.1021/ja1096682.

37. Sheridan KJ, Dolan SK, Doyle S. 2015. Endogenous cross-talk of fungal metabolites. Front Microbiol 5:732. https://doi.org/10.3389/fmicb.2014.00717.

38. Martens TM, Nunez O, Gallart-Ayala H, Leitao MC, Silva Pereira C. 2015. Plasma membrane permeabilisation by ionic liquids: a matter of charge. Green Chem 17:4587–4598. https://doi.org/10.1039/C5GC01472G.

39. Lange M, Muller C, Peiter E. 2014. Membrane-assisted culture of fungal mycelium on agar plates for RNA extraction and pharmacological analyses. Anal Biochem 453:58–60. https://doi.org/10.1016/j.ab.2014.02.023.

40. Petersen LM, Holm DK, Gotfredsen CH, Mortensen UH, Larsen TO. 2015. Investigation of a 6-MSA synthase gene cluster in "Aspergillus aculeatus" reveals 6-MSA-derived acyclic acid, acyclic A-8 and Epi-aculin A. ChemBiochem 16:2200–2204. https://doi.org/10.1002/cbic.201500210.

41. Rocheleau H, Al-Harthi R, Ouellert T. 2019. Degradation of salicylic acid by Fusarium graminearum. Fungal Biol 123:77–96. https://doi.org/10.1016/j.funbio.2018.11.002.

42. Siettmann R, Uebe R, Boer E, Bode R, Kunze G, Schauer F. 2010. Novel metabolic routes during the oxidation of hydroxylated aromatic acids by the yeast "Arxula adeninivorans." J Appl Microbiol 108:789–799. https://doi.org/10.1111/j.1365-2672.2009.04474.x.

43. Badawy AA. 2017. Kynurenine pathway of tryptophan metabolism: regulatory and functional aspects. J Int Tryptophan Res 10:2017846917691938. https://doi.org/10.1177/1178646917691938.

44. Heijman MJC, Wauben M, Welton JL, Zhao K, Mathivanan S. 2017. A comparative transcriptome analysis shows conserved metabolic regulation of a common precursor from two distinct genes generates metabolite diversity. Chem Sci 6:5913–5921. https://doi.org/10.1039/C6SC04910E.

45. Ye J, Coulouris G, Zaretskaya I, Cutcutache I, Rozen S, Madden TL. 2012. Primer-BLAST: a tool to design target-specific primers for polymerase chain reaction. BMC Bioinform 13:174. https://doi.org/10.1186/1471-2105-13-174.

46. Hartmann DO, Shumitz K, Siopa F, Leitao MC, Fiehn O, Downs DM, Hanson AD. 2015. Genomic and experimental validation of its human homologue in the yeast Candida albicans. PLoS One 8:e71364. https://doi.org/10.1371/journal.pone.0071364.
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the gene cluster level with MultiGeneBlast. Mol Biol Evol 30:1218–1223. https://doi.org/10.1093/molbev/msz025.

57. Blin K, Shaw S, Steinke K, Villebro R, Ziemert N, Lee SY, Medema MH, Weber T. 2019. antiSMASH 5.0: updates to the secondary metabolite genome mining pipeline. Nucleic Acids Res 47:W81–W87. https://doi.org/10.1093/nar/gkz310.

58. Grigoriev IV, Nikitin R, Haridas S, Kuo A, Ohm R, Otillar R, Riley R, Salamov A, Zhao X, Korzeniewski F, Smirnova T, Nordberg H, Dubchak I, Shabalov I. 2014. MycoCosm portal: gearing up for 1000 fungal genomes. Nucleic Acids Res 42:D699–D704. https://doi.org/10.1093/nar/gkt1183.

59. Khaldi N, Seifuuddin FT, Turner G, Haft D, Nierman WC, Wolfe KH, Fedorova ND. 2010. SMURF: genomic mapping of fungal secondary metabolite clusters. Fungal Genet Biol 47:736–741. https://doi.org/10.1016/j.fgb.2010.06.003.