Identification of a Conserved Transcriptional Activator-Repressor Module Controlling the Expression of Genes Involved in Tannic Acid Degradation and Gallic Acid Utilization in Aspergillus niger

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Tannic acid, a hydrolysable gallotannin present in plant tissues, consists of a central glucose molecule esterified with gallic acid molecules. Some microorganisms, including several Aspergillus species, can metabolize tannic acid by releasing gallic acid residues from tannic acid by secreting tannic acid specific esterases into the medium. The expression of these so-called tannases is induced by tannic acid or gallic acid. In this study, we identified a conserved transcriptional activator-repressor module involved in the regulation of predicted tannases and other genes involved in gallic acid metabolism.

The transcriptional activator-repressor module regulating tannic acid utilization resembles the transcriptional activator-repressor modules regulating galacturonic acid and quinic acid utilization. Like these modules, the Zn(II)₂Cys₆ transcriptional activator (TanR) and the putative repressor (TanX) are located adjacent to each other. Deletion of the transcriptional activator (ΔtanR) results in inability to grow on gallic acid and severely reduces growth on tannic acid. Deletion of the putative repressor gene (ΔtanX) results in the constitutive expression of tannases as well as other genes with mostly unknown function. Known microbial catabolic pathways for gallic acid utilization involve so-called ring cleavage enzymes, and two of these ring cleavage enzymes show increased expression in the ΔtanX mutant. However, deletion of these two genes, and even deletion of all 17 genes encoding potential ring cleavage enzymes, did not result in a gallic acid non-utilizing phenotype. Therefore, in A. niger gallic acid utilization involves a hitherto unknown pathway. Transcriptome analysis of the ΔtanX mutant identified several genes and gene clusters that were significantly induced compared to the parental strain. The involvement of a selection of these genes and gene clusters in gallic acid utilization was examined by constructing gene deletion mutants and testing their ability to grow on gallic acid. Only the deletion of a gene encoding an FAD-dependent monooxygenase...
INTRODUCTION

Tannins are polyphenolic aromatic compounds present in plant tissues such as leaves, bark, and wood (Aguilar et al., 2007). In plants, tannins are able to form strong complexes with proteins and polysaccharides like starch and cellulose to protect plants from degradation by fungal enzymes (Lekha and Lonsane, 1997). Tannic acid is a hydrolysable gallotannin, and consists of a central glucose molecule esterified with on average eight gallic acid molecules (Chávez-González et al., 2012).

Some microorganisms can use tannic acid as a carbon source. To utilize tannic acid, these microorganisms secrete tannin acyl hydrolases (also called tannases). Tannases catalyze the hydrolysis of the carboxylic ester bonds, resulting in the degradation of tannic acid to release gallic acid. Microorganisms that are capable of producing tannases include the bacterium Pseudomonas putida (Nogales et al., 2011), the yeast Blastobotrys (Arxula) adeninivorans (Meier et al., 2017) as well as several filamentous fungi of which Aspergillus species have been studied the most (Pinto et al., 2001; van Diepeningen et al., 2004). Although the amino acid sequences of bacterial, yeast, and fungal tannases are rather divergent, they all contain the active site motif of Gly-X-Ser-X-Gly, which is a characteristic of serine hydrolases (de las Rivas et al., 2019). The expression of fungal tannases is transcriptionally controlled. Their expression is induced when tannins or some of their hydrolysis products—such as gallic acid, pyrogallol, or methyl gallate—are present (Baipai and Patil, 1997; Lekha and Lonsane, 1997; Aguilar et al., 2007). However, gallic acid was also reported to repress tannase synthesis under solid state fermentation conditions in Aspergillus niger (Aguilar et al., 2001).

Based on available data, mainly derived from studies in bacteria, the ring structure in aromatic compounds, including gallic acid, is considered to be opened by specific so-called ring cleavage enzymes. Up to now, seven major monocyclic aromatics have been shown to be substrates of specific ring cleavage enzymes [reviewed by Lubbers et al. (2019b)]. These substrates, besides gallic acid, include catechol, protocatechuic acid, hydroxyquinol, hydroquinone, gentisic acid, and pyrogallol. After oxygenation, the ring-opened compounds are further metabolized and channeled into central metabolism (Lubbers et al., 2019b).

Two mechanisms of ring cleavage have been described and are catalyzed by either intradiol or extradiol ring cleavage dioxygenases. Ring cleavage using the intradiol dioxygenases, also called ortho-cleavage dioxygenases, takes place between hydroxyl groups of the ring structure, while the extradiol- or meta-cleavage dioxygenases cleave adjacent to a hydroxyl group (Harwood and Parales, 1996). Bacterial species use both intradiol and extradiol dioxygenases for the catabolism of aromatic compounds, but in fungi only intradiol cleavage has been described (e.g., Cain et al., 1968). In A. niger, four aromatic intradiol ring cleavage dioxygenases have been characterized for the catabolism of hydroxyquinol, catechol, and protocatechuate (Semana and Powlowski, 2019). The involvement of certain ring opening enzymes involved in protocatechuate was further confirmed by deletion analysis (Lubbers et al., 2019c). In the research presented here, we have identified several additional genes annotated as putative extradiol dioxygenase in the genome of A. niger whose encoding enzymes have not yet been studied.

The catabolic pathway of gallic acid in bacteria and yeast species has been studied in detail and shown in Figure 1. In P. putida, ring cleavage of gallic acid is catalyzed by the Fe^{2+}-dependent extradiol dioxygenase GaA, resulting in the intermediate 4-oxalomesaconic acid (OMA)keto, which is converted into OMAenol by the GaD isomerase. After hydration of OMAenol by the Zn^{2+}-dependent hydratase GalB, the degradation of the intermediate 4-carboxy-4-hydroxy-2-oxoacidic acid is catalyzed by GalC into pyruvic acid and oxaloacetic acid which channel into central metabolism (Nogales et al., 2011). A different metabolic pathway has been identified for the yeast B. adeniivorans. In this pathway, gallic acid is decarboxylated to form pyrogallol by the gallic acid decarboxylase Agdc1p. Ring cleavage of pyrogallol is believed to be catalyzed by a still unidentified catechol 1,2 dioxygenase to produce 2-hydroxymuconic acid, which is further degraded into pyruvate and acetaldehyde (Meier et al., 2017). Apart from Agdc1p, no other genes or gene products involved in the catabolism of gallic acid in B. adeniivorans have been identified.

Knowledge about the degradation pathways of aromatic acids in filamentous fungi is limited. Best studied is the 3-oxoadipate pathway, or β-ketoacid pathway, which has branches for the catabolism of protocatechuate and catechol using intradiol cleavage and channeling the products via β-ketoacid into the TCA cycle (Martins et al., 2015; Lubbers et al., 2019c, 2021). As indicated above, the intradiol dioxygenases involved have been characterized by Semana and Powlowski (2019) and Lubbers et al. (2019c). Additional enzymes involved in catabolic steps after ring cleavage of protocatechuic and salicylic acid have been identified in A. nidulans by proteomic, transcriptomics and deletion analyses (Martins et al., 2015) and in A. niger (Lubbers et al., 2019c, 2021). The catabolic pathway for cinnamic acid in A. niger has also been studied. Three enzymes play a role in cinnamic acid and sorbic acid degradation, but are not required for the catabolism of...
other related aromatic compounds (Lubbers et al., 2019a). Genes involved in the metabolism of gallic acid in filamentous fungi have yet not been reported.

Biochemical studies on the catabolism of gallic acid have been performed in A. niger and A. flavus (Watanabe, 1965; Gurujeyalakshmi and Mahadevan, 1987). Catabolic pathways for gallic acid were proposed based on the identification of intermediates. For A. niger, it is proposed that gallic acid is oxidized to a peroxide and decomposed to cis-aconitic acid and trans-aconitic acid, which can be decarboxylated to α-ketoglutaric acid (Watanabe, 1965). In A. flavus, the intermediates 4-carboxy-2-hydroxy-cis, cis-muconic acid (4-oxalomesaconate) and pyruvic acid were detected during gallic acid degradation, which suggests the involvement of an extradiol gallate dioxygenase enzyme (Gurujeyalakshmi and Mahadevan, 1987).

Saprophytic fungi, including A. niger, produce an arsenal of extracellular enzymes to degrade plant derived biomass (Pel et al., 2007; de Vries et al., 2017). The enzymes involved in the utilization and catabolism of plant compounds by fungi is tightly controlled. Transcription factors activate the expression of target genes, involved in specific metabolic pathways, by binding to specific transcription factor binding sites located upstream of these genes (Benocci et al., 2017). With such a controlled regulation system, transcriptional activation of substrate specific enzymes, transporters and catabolic pathway enzymes is only activated when a particular plant compound is present. Well-studied examples in A. niger are the transcription factors AmyR and XlnR that control the expression of genes encoding enzymes involved in starch and xylan utilization, respectively (van Peij et al., 1998; Petersen et al., 1999; van Kuyk et al., 2012). Recently, the regulatory mechanism of genes involved in the utilization of pectin and its main component galacturonic acid was studied (Alazi et al., 2016, 2017; Niu et al., 2017). These studies indicate that the galacturonic acid catabolic pathway in A. niger is controlled by a Zn(II)$_2$Cys$_6$ transcriptional activator (GaaR) in combination with a specific repressor protein (GaaX) (Niu et al., 2017). The activator (GaaR) and the corresponding repressor (GaaX) are located next to each other in the genome and form a module controlling the expression of genes involved in galacturonic acid utilization. Under non-inducing conditions, GaaX is expected to repress GaaR activity, thereby blocking the expression of target genes involved in the utilization of pectin and catabolism of galacturonic acid. A similar activator-repressor module for the degradation of quinic acid consisting of the Zn(II)$_2$Cys$_6$ transcriptional activator QutA/QA-1F and the corresponding repressor QutR/QA-1S is found in A. nidulans and Neurospora crassa, respectively (Giles et al., 1985; Geever et al., 1989; Levesley et al., 1996). When quinic acid or quinic acid derived metabolite binds to the repressor, it is expected...
TABLE 1 | Strains used in this study.

| Strain   | Genotype                                      | References          |
|----------|-----------------------------------------------|---------------------|
| NA02     | caspA                                         | Bos et al., 1988    |
| MA169.4  | caspA, pyrG378, kusA::DR-amdS-DR             | Carvalho et al., 2010|
| MA234.1  | caspA, kusA::DR-amdS-DR                      | Alazi et al., 2016  |
| MA514.1  | tanR::hygR in MA234.1                        | This study          |
| MA586.1  | tanX::hygR in MA234.1                        | This study          |
| TS6.4    | NRR3_04659::AopyrG in MA169.4                | This study          |
| MA831.1  | NRR3_06002::AopyrG in MA169.4                | This study          |
| TS2.1    | NRR3_06032::AopyrG in MA169.4                | This study          |
| MA835.1  | NRR3_04277::hygR in TS2.1                    | This study          |
| MA829.1  | NRR3_01651-01656::AopyrG                     | This study          |
| MA852.1  | NRR3_02180-02185::AopyrG                     | This study          |
| TS10.1   | NRR3_06147-06150::AopyrG                     | This study          |
| MA857.2  | NRR3_08290-08292::AopyrG                     | This study          |
| MA963.4  | NRR3_09054-09057::AopyrG                     | This study          |
| MA860.2  | NRR3_10365-10375::AopyrG                     | This study          |
| MA925.2  | ΔNRR3_00414, ΔNRR3_00984, ΔNRR3_01405, ΔNRR3_02496, ΔNRR3_02522, ΔNRR3_02644, ΔNRR3_02896, ΔNRR3_03141, ΔNRR3_03148, ΔNRR3_03326, ΔNRR3_04277, ΔNRR3_04335, ΔNRR3_04787, ΔNRR3_05330, ΔNRR3_05564, ΔNRR3_06632, ΔNRR3_11743 in MA234.1 | This study          |

This study

In this study, key players in the regulatory mechanism related to tannic acid utilization and gallic acid catabolism have been identified in A. niger. Its transcriptional control is mediated by an activator TanR and the corresponding putative repressor TanX. Like the GaaR/GaaX and QutA/QutR activator-repressor modules, TanR [Zn(II)2Cy6 activator] and TanX (repressor) are located next to each other in the genome.

MATERIALS AND METHODS

Strains, Media, and Growth Conditions
The A. niger strains used in this study are listed in Table 1. Strains were grown on solid or liquid medium [containing 1.5% (w/v) Scharlau agar] minimal medium (MM) or on complete medium (CM) as described (Arentshorst et al., 2012). Aspergillus niger transformants were purified as described (Arentshorst et al., 2012) using a final concentration of 100 µg/mL hygromycin. Mycelial growth assays were performed on MM plates, usually supplemented with 5 mM of the aromatic compound. Aromatic compounds were weighed, dissolved in sterile water and mixed with an equal volume of 2 x concentrated solidified MM. Radial growth of A. niger strains was assayed by point inoculation of 5 µL filtered spore suspension (2*10^5 spores/mL) in the center of an agar plate and incubation of the plates for 5 days at 30°C. Escherichia coli DH5α was used for plasmid construction and cultured at 37°C in Luria-Bertani medium, with ampicillin (100 µg/mL).

Bioreactor Cultures, RNA Isolation, and RNAseq
Bioreactor controlled batch fermentation on fructose of the ∆tanX strain MA586.1 was performed as described previously (Jørgensen et al., 2010; Niu et al., 2017). In short, autoclaved bioreactor vessels were filled with 5 L of sterile MM containing 0.75% fructose as a carbon source. During cultivation at 30°C, the controller was set to maintain pH 3 by addition of titrants (2 M NaOH or 1 M HCl). Sterile air was supplied at a rate of 1 L min⁻¹. Prior to inoculation, 1.5 ml of 10% (w/v) filter-sterilized yeast extract was added to enhance conidial germination. Cultures were inoculated with freshly harvested spores at a concentration of 7.0 × 10⁴ conidia per liter. To reduce the loss of hydrophobic conidia during germination, the stirrer speed was set to 250 rpm and the culture was aerated via the headspace during the first 6 h after inoculation. Subsequently, the stirrer speed was increased to 750 rpm, 0.5 ml of polypropylene glycol P2000 was added as an antifoam agent and air was supplied via the sparger. Cultures broth was harvested at regular intervals from batch cultures and mycelial biomass was retained by vacuum filtration using glass microfiber filters (Whatman). Both biomass and filtrate were quickly frozen in liquid nitrogen and subsequently stored at −80°C. Dry biomass concentrations were gravimetrically determined from lyophilized mycelia originating from a known mass of culture broth. RNA was isolated from mycelium that was grown until mid-exponential phase. Extraction of RNA, library construction and RNA sequencing were performed as described (Niu et al., 2017). Three biological replicates for the reference strain and two biological replicates for the ∆tanX strain were performed. For each RNAseq sample a dataset of at least 1.5 Gb (20 million reads) of Illumina-filtered sequence data was obtained. Processing of the RNA seq data and differences in transcript expression between genotypes were analyzed with DESeq2 as described (Niu et al., 2017 and references therein).

Molecular Techniques and Construction of Gene Deletion Strains
The gene deletion strains used in this study were created using either the split marker method (Arentshorst et al., 2015), or by using the marker-free CRISPR/Cas9 genome editing method (van Leeuwe et al., 2019). For PCR amplifications Phire Hot Start II DNA Polymerase (Thermo Scientific) was used. Amplification and purification of split marker fragments were done as described (Arentshorst et al., 2015) using gene specific primer sets which are listed in Supplementary Table 1. The tanX and tanR genes were deleted using the split marker method using hygromycin as selection marker. Deletion of tanX was confirmed by Southern blot analysis (Supplementary Figure 1) and deletion of tanR was verified via diagnostic PCR (Supplementary Figure 2). Deletion of NRR3_06002 was performed with split marker fragments containing the Aspergillus
oryzae pyrG marker (AopyrG) and putative deletions strain were analyzed via diagnostic PCR (Supplementary Figure 3).

Deletion of the six gene clusters that were induced in the ΔtanX mutant (ΔNRRL3_01651-NRRL3_01656, ΔNRRL3_02180-NRRL3_02185, ΔNRRL3_06147-NRRL3_06150, ΔNRRL3_08290-NRRL3_08292, ΔNRRL3_09054-NRRL3_09057, and ΔNRRL3_10365-NRRL3_10375) was performed with split marker fragments and putative deletions strain were verified by diagnostic PCR (Supplementary Figures 4–9). Single deletion of NRRL3_08632 and double deletion of NRRL3_08632 and NRRL3_04277 were made by split markers using the AopyrG marker and hygromycin marker, respectively (Supplementary Figure 10). Deletion of NRRL3_04659 was performed with split marker fragments and AopyrG and proper deletion was verified by diagnostic PCR (Supplementary Figure 11).

To create a strain with all the seventeen putative ring cleavage enzymes deleted (Δrce 17x) from its genome, the marker-free CRISPR/Cas9 genome editing method (van Leeuwe et al., 2019) was used. To increase the chance of successfully creating a double strand break in the gene of interest, two sgRNA targets per gene were designed using the CHOPCHOP webtool (Labun et al., 2019). For each gene two pFC332-based Cas9-hygR-AMA1 vectors each containing a unique Pro1-sgRNA expression cassette were constructed. The guide RNAs were PCR amplified (Supplementary Table 1) and cloned into pFC332. The sequence of final plasmids was verified by sequencing (Macrogen, The Netherlands).

For each gene deletion, a repair DNA fragment was constructed to allow homologous recombination at the locus of the gene of interest. This repair DNA fragment consists of the 5′ flank and the 3′ flank of the gene, separated by a 23 bp spacer. The flanks of all ring cleavage enzymes were PCR amplified with genomic DNA of N402 as template and primers that are listed in Supplementary Table 1. The overlapping spacer sequence in all PCR products is used in a subsequent fusion PCR to fuse together the 5′ flank and 3′ flank, resulting in the repair DNA fragment. The sizes of the flanks and repair DNA fragments for each ring cleavage enzyme are shown in Supplementary Table 1. Deletion of genes encoding the ring cleavage enzymes was achieved by transforming strain MA234.1 (Δaku70) with plasmids pFC332-gRNA1 and pFC332-gRNA2 (both 2–5 μg DNA) and the repair DNA fragment (1–2 μg DNA). Transformants were purified on MM supplemented with hygromycin, followed by a purification step on MM to allow loss of the hygR-AMA1 pFC332 plasmids. Afterwards the loss of these plasmids was confirmed by growth analyses on MM supplemented with hygromycin and correct integration of the repair DNA was verified by diagnostic PCR. In one transformation two or three genes were deleted simultaneously resulting in the seventeen ring-cleavage enzyme deletion strain MA925.2 (Table 1), confirmed by diagnostic PCR (Supplementary Figure 12).

**Metabolite Analysis**

The extracellular metabolite production by parental and mutant *A. niger* strains was performed using a two-step culture; a primary culture generating fungal biomass and a secondary culture to induce protein and metabolite production. For primary culture generation, 100 mL of CM with 2% fructose was inoculated with 2 × 10⁹ spores/mL and incubated for 24 h at 30°C with shaking at 220 rpm. Primary cultures were filtered through with a Buchner funnel lined with Miracloth (Calbiochem). The collected mycelia was rinsed with sterile ddH₂O and a volume of 5 mL of mycelia was used to start secondary cultures in 250 mL flasks containing 50 mL MM, 1% gallic acid, and 0.1% fructose. Secondary cultures were incubated at 30°C with shaking at 220 rpm. Two mL of culture liquid was harvested at 0, 4, 8, 18, and 24 h. For each time point, the sample was centrifuged at 15 000 × g for 15 min. A 100 μL aliquot of supernatant was transferred to a tube with 200 μL cold methanol and left at −20°C overnight. The samples were centrifuged at 15 000 × g for 30 min, then a 30 μL aliquot of the supernatant was mixed with 150 μL of 0.1% formic acid. A 10 μL aliquot was analyzed by high resolution liquid chromatography-mass spectrometry using an Agilent 1,260 Infinity II HPLC system (Agilent technologies, Santa Clara, CA, USA) connected in-line to 7 Tesla Thermo-Finnigan LTQ-FT mass spectrometer (Thermo Electron Corporation, San Jose, CA, USA). Chromatography separation of sample components was done using a Synergi Hydro-RP 150 × 2.0 mm, 4 μm column (Phenomenex, Torrence, CA, USA). The solvents used to generate the gradient during reversed-phase separation were 0.1% formic acid in water for Solvent A and 0.1% formic acid in acetonitrile for Solvent B. Solvent flow rate was 300 μL/min and the gradient started at 1% B, increased to 30% B in 6 min, increased to 80% B in 2 min, increased to 90% B in 0.1 min, maintained at 90% for 1 min, decreased to 1% B in 0.1 min and kept at 1% B for 4 min. Column eluate was connected to a Thermo-Finnigan Ion Max electrospray source. Spectra were acquired in negative mode from 60 to 800 m/z at 50,000 resolution at 200 m/z. LC-MS data were processed for ion feature extraction and compound annotation using Compound Discoverer 3.0 (Thermo-Fisher Scientific, Waltham, MA). Compound annotation was done by querying the metabolite databases from ChEBI, ChEMBL, and MoNA using a 5 ppm mass accuracy search window criteria. The relative abundance of selected compounds of interest were compared across samples using extracted chromatogram peak area values calculated by the Xcalibur 2.2. Quan Browser software (Thermo-Fisher Scientific, Waltham, MA).

**RESULTS**

**Tannases Are Constitutively Expressed in Putative Repressor Mutant ΔNRRL3_08276**

We identified a transcriptional activator-repressor module in *A. niger* which shows similarities in both genomic organization and protein sequence to the activator-repressor modules involved in galacturonic acid or quinic acid utilization (Niu et al., 2017; Figure 2A). The transcriptional activator [NRRL3_08275 (An03g6810)] is a Zn(II)₂Cys₆ type transcription factor. The predicted repressor [NRRL3_08276 (An03g06800)] shows homology to repressor proteins that are involved in galacturonic acid ([NRRL3_08194 (GaaX)] and quinic acid

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FIGURE 2 | Schematic overview of the genomic organization of activator/repressor modules in the genome of A. niger and the presence of PFAM domains in putative repressor proteins. (A) Four modules consisting of an activator and a putative repressor protein are present in the genome of A. niger. The genes encoding the activator (Continued)
FIGURE 2 | and repressor proteins are localized adjacent to each other and either expressed in a head to tail orientation (gaaR/gaaX) or a potential bidirectional promoter (qutA/qutR; tanR/tanX) and NRRL3_07604/NRRL3_07605). The length of the bidirectional promoter is indicated. (B) Schematic representation of the phylogeny and relevant PFAM present in identified putative repressor protein in A. niger (NRRL3), A. nidulans (AN), and N. crassa (NcU), compared to AROM. Protein sequences were retrieved using BLASTP with the AromA protein sequence of A. niger (NRRL3_11273) as a query against A. nidulans (FGSCA4) or N. crassa (OR74A). Orthologs of identified repressor in the tree fungal species are indicated by the colored boxes. PFAM searches were performed at https://www.ebi.ac.uk/Tools/pfa/pfamscan/. The unrooted tree was constructed using standard settings in Clustal Omega at https://www.ebi.ac.uk/Tools/msa/clustalo/.

(NRRL3_11039 (QutR)) utilization and a fourth putative repressor (NRRL3_07605) with unknown function (Niu et al., 2017). The family of four putative repressor proteins in A. niger identified previously (Niu et al., 2017) consists of proteins with conserved protein family domains (PFAM domains) that are similarly organized as in the AROM (aromatic biosynthesis) protein. AROM is a conserved and large protein (1,586 amino acids long in A. niger) composed of five domains, and the individual domains are required for five enzymatic steps representing the prechorismate shikimate pathway which is required for aromatic amino acid biosynthesis (Duncan et al., 1987). A BLASTP search using AROM1 protein of A. niger (NRRL3_11273) as a query identified the four putative repressor proteins in A. niger. The homology between AROM and the putative repressor is found in the last three domains of the AROM protein. These last three domains of AROM consist of a shikimate kinase domain (PF01202.22), a 3-dehydroquininate dehydratase domain (PF01487.15) and a shikimate dehydrogenase domain (PF08501.11 and PF18317.1). These domains are also present in the orthologous repressor protein identified in A. nidulans and N. crassa (Figure 2B).

The putative repressor protein (NRRL3_08194) was found in BLASTP searches using AROM as a query, but the PF1202.22, PF8501.11, and PF18317.15 domains in the GaaX paralogs are not recognized in PFAM searches (Figure 2B). In addition, PF01487.15 was not detected in NRRL3_07605.

Loss of function of the galacturonic acid specific repressor protein (GaaX) or the quinic acid specific repressor protein (QutR) results in constitutive activation of expression of genes involved in pectin utilization and galacturonic acid metabolism (ΔgaaX) or quinic acid metabolism (ΔqutR), respectively (Niu et al., 2017; Ram and Tsang, unpublished data). To identify metabolic pathways which could be controlled by the NRRL3_08275-NRRL3_08276 module, a RNAseq study with deletion mutant of the putative repressor gene [MA586.1 (ΔNRRL3_08276)] was performed (see Materials and Methods for details and Supplementary Figure 1 for strain verification). No growth abnormality in terms of radial growth or sporulation was observed for the deletion mutant compared to the parental strain when cultured on minimal medium agar plates with glucose as carbon source (data not shown).

To identify which metabolic pathways are controlled by this regulatory module, RNAseq analysis was carried out. The deletion mutant (ΔNRRL3_08276) and its parent (MA234.1), were cultivated in bioreactors and RNA was isolated from exponentially growing cells on fructose medium. Fructose was chosen as a carbon source as it is generally less strong as a carbon catabolite repressing sugar compared to glucose. The specific growth rate (0.217 ± 0.002 g dry weight/kg broth/h) and maximal biomass accumulation (4.289 ± 0.194 g dry weight/kg broth) of the ΔNRRL3_08276 strain were similar to the growth rate and maximal biomass accumulation (0.214 ± 0.007 g dry weight/kg broth/h and 4.151 ± 0.134 g dry weight/kg broth) of parental strain MA234.1. The cultures and transcriptomic data of MA234.1 used for comparison have been previously described (Niu et al., 2017). Total RNA was isolated from A. niger strain ΔNRRL3_08276 and MA234.1 when 70% of the maximal biomass was reached. RNAseq analysis was performed to identify genes that were significantly higher expressed in the ΔNRRL3_08276 mutant than in the parental strain (Supplementary Table 2). A DESeq2 analysis on the transcriptome data was performed to define differentially expressed genes. This analysis indicated that 330 genes displayed significantly higher expression (FC > 2 and FDR < 0.001) (Supplementary Table 2). Among the twenty genes with the largest expression increase in the putative repressor mutant compared to the parental strain (Table 2) are two genes (NRRL3_00063, NRRL3_06367) encoding predicted tannases, suggesting that the NRRL3_08275/NRRL3_08276 module regulates genes involved in tannic acid utilization.

The Zn(II)2Cys6 Transcription Factor NRRL3_08275 Is Required for Growth on Tannic Acid and Gallic Acid

To test whether the module is indeed related to tannic acid utilization, the Zn(II)2Cys6 transcriptional activator NRRL3_08275 (An0366810) was deleted (Supplementary Figure 2). Growth on substrates including tannic acid and its main monomeric constituent gallic acid was analyzed. As shown in Figure 3, growth of ΔNRRL3_08275 was severely reduced on tannic acid and no growth occurred on gallic acid, confirming that NRRL3_08275 regulates tannic acid degradation. To determine whether the growth defect of ΔNRRL3_08275 was specific for tannic acid and gallic acid, growth of ΔNRRL3_08275 on galacturonic acid, quinic acid (Figure 3) and on various aromatics including ferulic acid, p-hydroxybenzoic acid, protocatechuic acid, caffeic acid, vanillic acid, p-coumaric acid, gentisic (2,5-dihydroxybenzoic) acid, salicylic acid, and catechol (Supplementary Figure 13) was tested. ΔNRRL3_08275 grew on these substrates equally well as the control strain (MA234.1), indicating that the growth defect of ΔNRRL3_08275 is specifically related to the utilization of tannic acid and its main degradation product gallic acid (Figure 3; Supplementary Figure 13). Growth of the putative repressor knockout (ΔNRRL3_08276) was not affected by any of the compounds tested (Figure 3; Supplementary Figure 13). We therefore refer to the Zn(II)2Cys6 transcriptional activator.
TABLE 2 | Top 20 of genes with the highest fold change (FC) in expression between the ΔtanX mutant and the parental strain during exponential growth on fructose.

| Transcript | Description | FC (Dseq²) | FDR |
|------------|-------------|------------|-----|
| NRRL3_01655* | O-methyltransferase, COMT-type | 5,712.8 | 0 |
| NRRL3_04659 | FAD-binding protein, putative monoxygenase | 2,392.2 | 0 |
| NRRL3_03155 | Ser-Thr-rich GPI-anchored protein | 452.7 | 0 |
| NRRL3_01653* | Multicopper oxidase | 297.1 | 3.37E⁻¹⁵⁰ |
| NRRL3_08632 | Extradiol ring-cleavage dioxygenase | 232.7 | 2.64E⁻²⁰⁴ |
| NRRL3_01658* | Hypothetical protein | 189.8 | 6.69E⁻¹⁶⁷ |
| NRRL3_03935 | MFS-type transporter | 179.7 | 0 |
| NRRL3_07468 | Carboxylesterase | 142.5 | 7.40E⁻²⁶⁸ |
| NRRL3_00063 | Tannase/feruloyl esterase family protein | 111.3 | 0 |
| NRRL3_06002 | Putative decarboxylase, SnoaL-like domain containing protein | 93.8 | 0 |
| NRRL3_01654* | Prenyltransferase, UbiA family | 94.5 | 3.87E⁻⁷⁷ |
| NRRL3_06149b | Hypothetical protein | 93.1 | 0 |
| NRRL3_09057* | PAN-1 domain-containing protein | 79.8 | 3.94E⁻⁶⁷ |
| NRRL3_10195 | WW domain-containing protein | 65.9 | 9.40E⁻²⁷⁶ |
| NRRL3_11834 | ABC transporter, family 2, CDR-type | 65.6 | 3.69E⁻⁶⁵ |
| NRRL3_09055* | Hypothetical protein | 65.0 | 1.32E⁻⁵⁸ |
| NRRL3_09054* | Glucan endo-1,3-alpha-glucosidase | 57.7 | 4.40E⁻⁶⁴ |
| NRRL3_04136 | Guanyl-Specific ribonuclease | 50.6 | 4.17E⁻²¹⁸ |
| NRRL3_08367 | Tannase/feruloyl esterase family protein | 47.6 | 1.59E⁻⁹⁰ |
| NRRL3_06150a | Isochorismatase family protein | 40.7 | 1.26E⁻⁵² |

FC, Fold change; FDR, False discovery rate.

* Genes belonging to gene cluster comprising NRRL3_01649-NRRL3_01658.

b Genes belonging to gene cluster comprising NRRL3_09054-NRRL3_09057.

FIGURE 3 | Growth analysis of A. niger reference strain (MA234.1), ΔtanR, ΔtanX, Δrce 17x, and ΔgacA (ΔNRRL3_04659) on various carbon sources. Spores were point inoculated on minimal medium (MM) supplemented with the indicated carbon source (5 mM). Strains were grown for 5 days at 30°C before pictures were taken.
NRRL3_08275 as TanR and the putative repressor protein NRRL3_08276 as TanX, analogous to the GaaR/GaaX activator-repressor module (Niu et al., 2017). The expression of the tanR transcription factor is 4-fold induced in the ΔtanX mutant (Supplementary Table 2). This upregulation of the tanR transcription factor in the absence of the putative repressor suggests that tanR itself is also controlled by the tanR/tanX module and tanR can upregulate itself.

A Putative Gallic Acid Decarboxylase That Is Induced in the ΔtanX Strain Is Not Required for Growth on Gallic Acid

Having established that the TanR/TanX module is likely to control degradation of tannic acid by inducing the expression of predicted tannases and the utilization of gallic acid, we further examined the gene expression pattern of the tanX mutant compared to its parental strain for genes possibly involved in catabolism of gallic acid (Table 2). The first candidate gene was identified based on homology to Agdc1p, a gallic acid decarboxylase involved in the utilization of gallic acid in the yeast B. adeninivorans. BLASTp searches using Agdc1p identified a possible ortholog, NRRL3_06002, which showed the highest similarity (51% identity and 69% similarity) to Agdc1p. The expression of NRRL3_06002 is strongly induced in the ΔtanX mutant compared to the parental strain (FC = 95.9, FDR < 0.001) (Table 2, Supplementary Table 2). A deletion mutant of NRRL3_06002 was constructed, verified via diagnostic PCR (Supplementary Figure 3), and its growth on tannic acid and gallic acid analyzed. The ΔNRRL3_06002 mutant grew normally on tannic acid and gallic acid, indicating that this gene is not required for growth on gallic acid. This result therefore indicates that gallic acid catabolism in A. niger is different from that of B. adeninivorans, possibly via redundant or alternative pathways.

Gene Clusters That Are Upregulated in the ΔtanX Strain Are Dispensable for Growth on Gallic Acid

Six gene clusters (NRRL3_01651-NRRL3_01656; NRRL3_02180-NRRL3_02185; NRRL3_06147-NRRL3_06150; NRRL3_08290-NRRL3_08292; NRRL3_09054-NRRL3_09057; NRRL3_10365-NRRL3_10375) were identified because the genes in these clusters were upregulated in the ΔtanX mutant compared to the parental strain (Supplementary Table 3). Since in some cases genes encoding enzymes involved in catabolic pathways are clustered (e.g., quinic acid or galactarconic acid), we examined the possibility that genes in these clusters were involved in tannic acid or gallic acid utilization by deleting these gene clusters. Deletion mutants [ΔNRRL3_01651-NRRL3_01656 (MA829.1); ΔNRRL3_02180-NRRL3_02185 (MA852.1); ΔNRRL3_06147-NRRL3_06150 (TS10.1); ΔNRRL3_08290-NRRL3_08292 (MA857.2); ΔNRRL3_09054-NRRL3_09057 (MA963.4); ΔNRRL3_10365-NRRL3_10375 (MA860.2)] were verified by diagnostic PCR (Supplementary Figures 4–9) and tested for growth on gallic acid and tannic acid. All deletion mutants grew normally on gallic acid and tannic acid indicating that these gene clusters are not exclusively required for tannic acid degradation and gallic acid utilization (data not shown). Since the gene clusters were deleted individually, it cannot be formally excluded that these gene clusters have redundant functions in gallic acid catabolism.

Intra- or Extradial Ring Cleavage Enzymes in the A. niger Genome Are Not Required for Growth on Gallic Acid

In a further attempt to identify genes encoding proteins involved in gallic acid utilization, the role of putative ring cleavage enzymes, identified to be relevant for growth on gallic acid in bacteria, was addressed. Two putative ring cleavage enzymes encoded by NRRL3_08632 and NRRL3_04277 were 232 and 38-fold induced in the ΔtanX mutant (Table 2, Supplementary Table 4). Single and double deletion mutants were constructed, verified (Supplementary Figure 10), and tested for growth on gallic acid and tannic acid. The growth on gallic acid of either single mutants or the double mutant was not affected (data not shown) indicating that the ring cleavage of gallic acid does not exclusively depend on these two ring cleavage enzymes, leaving the possibility that other ring cleavage enzymes might take over their function. To address this possibility, we searched the A. niger genome and identified seventeen genes encoding potential ring cleavage enzymes based on BlastP results in combination with Pfam domain analysis. These seventeen proteins were selected because of the presence of PF00775.11, PF02900.8, and/or PF04444.4 domains which are related to a dioxygenase_C domain, the catalytic LigB subunit of aromatic ringopening dioxygenase domain and/or the Dioxygenase_N domain, respectively. The domain structure in combination with the phylogenetic relationship of these proteins are shown in Figure 4. The expression of these seventeen genes was analyzed in the ΔtanX parental expression data set and is shown in Supplementary Table 4. To address the possibility that functionally redundant ring cleavage enzymes occur within this group of seventeen enzymes, we constructed an A. niger strain in which all seventeen genes encoding ring cleavage enzymes were deleted (Supplementary Figure 12). Growth of the 17-fold ring cleavage enzyme mutant on gallic acid or tannic acid was not impaired compared to the parental strain, indicating that none of the ring cleavage enzymes is required for gallic acid utilization (Figure 3). Growth of the 17-fold deletion strain on quinic acid is abolished (Figure 3) which is due to deletion of NRRL3_01405 (Sgro, Arentshorst, Ram and Tsang, manuscript in preparation).

A FAD Domain-Containing Monooxygenase Is Required for Growth on Gallic Acid

In the transcriptome data the second most induced gene (Table 2) is NRRL3_04659. This gene encodes a FAD domain-containing monooxygenase with sequence similarity to salicylate hydroxylase (NahG) of P. putida. As ring cleavage has also been reported to be mediated by monooxygenases and because of its strong induction in the ΔtanX mutant, a knockout strain for this gene was made and verified by diagnostic PCR (Supplementary Figure 11). Growth assays on various aromatic compounds revealed that the ΔNRRL3_04659 mutant
does not grow on gallic acid and shows strongly reduced growth on tannic acid (Figure 3). Growth on galacturonic acid, quinic acid, and aromatic compounds such as ferulic acid, p-hydroxybenzoic acid, protocatechuic acid, caffeic acid, vanillic acid, p-coumaric acid, gentisic acid, salicylic acid, and catechol was not affected by deletion of NRRL3_04659 (Figure 3; Supplementary Figure 13), implying that this monooxygenase gene, which we called gacA for gallic acid catabolism, is required and specific for the initial ring cleavage of gallic acid.

Growing the ΔgacA on agar plates containing 2.5 mM fructose as a carbon source and comparing the growth on agar plates containing 2.5 mM fructose and 25 mM gallic acid revealed that the addition of gallic acid reduced growth of the ΔgacA mutant (as well as the ΔtanR mutant) compared to the parental strain (Supplementary Figure 14). This preliminary result indicates that the ability to metabolize gallic acid also contributes to the detoxification of gallic acid.

Further indications that GacA is required for ring cleavage of gallic acid was obtained by extracellular metabolite analysis. The parental strain (MA234.1) and the ΔgacA strains (TS6.4) were grown on fructose/gallic acid medium for 18 and 24 h. The ΔtanR strain was also included in the same experiment. Extracellular metabolites present in the medium after 18 and 24 h were detected by LC-MS-MS as described in the Materials and Methods. The analysis showed that levels of gallic acid in the medium of the ΔgacA remain...
FIGURE 5 | (A) Extracellular metabolites analysis of A. niger strains grown after transfer to 1% gallic acid/0.1% fructose. A. niger strains MA234.1 (reference strain), the ΔtanR mutant, and the ΔgacA (NRRL3_04659) mutant were pregrown in CM supplemented with fructose for 24 h, mycelium was washed and transferred MM supplemented with 1% gallic acid/0.1% fructose for the indicated times. Most abundant extracellular metabolites (gallic acid, 2-carboxy-cis,cis-muconate and cis,cis-muconate) were detected and their relative abundance over time course (0, 4, 8, 16, and 24 h after transfer) were analyzed as described in Materials and Methods. (B) Proposed first steps in catabolism of gallic acid in A. niger based on the accumulation of intermediates in the ΔtanR mutant.

the same during the time course (Figure 5A). In contrast, gallic acid levels in the parental strain (MA234.1) dropped over time, and after 24 h of growth hardly any gallic acid remained. Interestingly, in the ΔtanR mutant, gallic acid levels dropped over time and other metabolites which could be identified as muconate and 2-carboxy-cis,cis-muconate were accumulating in the medium. Possibly 2-carboxy-cis,cis-muconate and muconate are intermediates in the catabolism of gallic acid and the expression of the gene encoding the enzyme acting on 2-carboxy-cis,cis-muconate is no longer induced in the ΔtanR mutant (Figure 5B). This could result in accumulation of 2-carboxy-cis-cis-muconate if gacA was still expressed to sufficient levels to convert gallic acid into 2-carboxy-cis-cis-muconate.

DISCUSSION

In the genome of A. niger over 440 different Zn(II)$_2$Cys$_6$ transcription factors can be found (Ram, unpublished). Most of the Zn(II)$_2$Cys$_6$ transcription factors are expected to be transcriptional activators and are responsible for the controlled expression of their target genes in response to the right inducer. The functional characterization of this large family of Zn(II)$_2$Cys$_6$ transcription factors is still very limited and even less is known about the detailed activation mechanisms. Among the large set of Zn(II)$_2$Cys$_6$ transcription factors in A. niger, we previously noticed the presence of four Zn(II)$_2$Cys$_6$ transcription factors, each of their encoding gene is co-localized in the genome with a gene encoding a putative repressor protein.
One of the Zn(II)2Cys6 transcription factors that is clustered with a repressor is the transcriptional activator involved in quinic acid utilization (QutA). This system has been studied in detail both in *A. nidulans* and in *N. crassa* (Giles et al., 1991). The repressor protein (QutR) shows significant sequence similarity toward the C-terminal half of the AROM protein (Figure 1B). AROM is a large multidomain protein and individual domains are involved in five enzymatic steps in the prechorismate pathway needed for the biosynthesis of aromatic amino acids (Duncan et al., 1987). The working hypothesis is that the repressor is a catalytically inactive protein that can still recognize or bind to the inducer. Apart from the activator-repressor module that regulates quinic acid utilization, a second module (Gaar/GaaX) has been functionally characterized; this module is involved in the regulation of genes involved in (poly)-galacturonic acid (pectin) degradation and utilization (Niu et al., 2017). In the current study, we show that a third activator and putative repressor module (TanR and TanX) is required for tannic acid degradation and gallic acid utilization. The function of the fourth activator-repressor module (NRRL3_07604 and NRRL3_07605) is currently unknown. All four putative repressor proteins show low sequence similarity toward each other and to the AROM1 protein. The mechanism by which the putative repressor and activator interact is unknown, but it has been suggested that an inducer molecule binds to the putative repressor protein which results in dissociation of the repressor from its conjugate transcriptional activator. The release of the repressor is suggested to result in an active transcription factor (Giles et al., 1991; Niu et al., 2017). The notion that all repressors show sequence similarity with AROM indicates that these putative repressors share a common evolutionary origin, and how these conserved modules have evolved is an intriguing question for further research.

The transcriptomic analysis of the putative repressor knockout mutant (ΔtanX) revealed a number of genes under the control of the TanR/TanX module. By analogy with the studies we carried out with the repressor mutant related to galacturonic acid utilization (ΔgaaA mutant), we expected target genes of the module to be expressed in higher levels in the repressor knockout mutant compared to the parental strain. In this study, our interest was drawn to the genes involved in the utilization of gallic acid which is not well-characterized in fungi. One of the most highly induced genes, NRRL3_06002, encodes a putative decarboxylase with high sequence similarity to a gallic acid decarboxylase from the yeast *B. adeninivorans* (Agdc1). In *B. adeninivorans*, gallic acid is degraded to pyrogallol before further degradation via 2-hydroxyxymuconic acid and oxalocrotonate (Meier et al., 2017). *A. niger* seems to have a different, or at least an alternative, catabolic pathway since the deletion of NRRL3_06002 did not result in a gallic acid non-utilizing phenotype. The possibility of redundancy of a gallic acid decarboxylase was examined and three homologs of NRRL3_06002 (NRRL3_06896, NRRL3_07601 and NRRL3_07467) were identified, but none of them showed a >2-fold higher induction in the ΔtanX mutant compared to the parental strain. The expression values of the three homologs were relatively low in either the reference strain or the ΔtanX mutant (Supplementary Table 2; Table 2), making it unlikely that one of these genes is responsible for taking over the function of NRRL3_06002. In agreement with that, the metabolite analysis in the parental strain, the ΔtanR mutant, and the ΔgaaA (ΔNRRL3_04659) mutant did not reveal the presence of pyrogallol, indicating that the degradation pathway of gallic acid does not involve direct decarboxylation from gallic acid to pyrogallol as an intermediate as in the case of *B. adeninivorans* (Meier et al., 2017), *A. oryzae* (Guo et al., 2014) and some bacterial species (Jiménez et al., 2013; Sonia et al., 2017).

Analysis of the top 20 genes that were fold-change wise the highest expressed in the ΔtanX strain compared to the parental strain identified several genes that were clustered to each other (Table 2). Therefore, the entire transcriptome data was analyzed for gene clusters that were significantly upregulated in the ΔtanX mutant compared to the parental strain. This resulted in the identification of six clusters consisting of three genes or more that were induced in the ΔtanX mutant (Supplementary Table 3). These gene clusters were individually deleted from the genome, but the deletion mutants were still able to grow as well as the parental strains on gallic acid indicating that none of the genes in these clusters is exclusively required for gallic acid utilization. Two gene clusters (cluster NRRL3_10365-NRRL3_10375 and NRRL3_02180-NRRL3_02185) contain a polyketide synthase encoding gene and a non-ribosomal peptide encoding gene (NRRL3_10375 and NRRL3_02185, respectively), which suggests that these clusters could be related to secondary metabolite production.

The molecular mechanism on how the putative TanX repressor protein controls the activity of the TanR transcription factor is not known. By analogy with the modules regulating quinic acid utilization in *A. nidulans* (QutA/QutR) and *N. crassa* (QA-1F/QA-1S) and galacturonic acid utilization in *A. niger* (Gaar/GaaX) a highly speculative model can be envisioned. In this model, we expect the transcriptional repressor and the transcriptional activator to be bound to each other under non-inducing conditions, leading to an inactive transcription factor complex. Upon the presence of an inducer, we expect binding of the inducer to the putative repressor protein as discussed previously (Niu et al., 2017), leading to the dissociation of the putative repressor from the transcription factor/repressor complex. Dissociation of the putative repressor from the transcription factor is expected to result in an active transcription factor. For quinic acid utilization, quinic acid or a metabolic derivative from quinic acid (Huiet and Giles, 1986) and in the case of galacturonic acid utilization 2-keto-3-deoxylgalactonate (Alazi et al., 2017) are indicated to be this inducer, while for the gallic acid utilization pathway this inducer is currently unknown. In addition to regulation of the activity of TanR at the protein level, the TanR transcription factor is also transcriptionally regulated as we observed a 4-fold induction of tanR expression in the ΔtanX mutant, indicating a positive feedback loop in that TanR can also induce its own expression. It should be noted that the current model of interactions between the inducer, putative repressor and transcriptional activator are highly speculative and needs to be tested in future studies.

Another intriguing result of our study is that in *A. niger* potential ring cleavage enzymes (both intra- and extradiol acting) are not involved in gallic acid utilization. Even though two potential ring cleavage enzymes (NRRL3_04277...
and NRRL3_08632) were highly induced in the ΔtanX strain, their simultaneous deletion did not result in a growth defect on gallic acid. To rule out the possibility that other ring cleavage enzymes would take over each other’s function in gallic acid utilization, all seventeen genes annotated to encode potential ring cleavage enzymes were deleted using marker-free CRISPR-CAS9 genome editing. The 17-fold mutant was still able to grow on gallic acid, indicating an alternative way of ring-opening. The proposed function of two intradiol-acting dioxygenases based on their biochemical characterization (Semana and Powalkowski, 2019) was confirmed in the deletion strains. NRRL3_01405 (prcA) was predicted to encode a protocatechuate-3,4-dioxygenase and deletion of NRRL3_01405 resulted in the inability to grow on protocatechuc acid (Lubbers et al., 2019; Sgro et al., unpublished), explaining the inability of the 17-fold deletion mutant to grow on quinic acid (Figure 3).

Likewise, NRRL3_04787 was predicted to encode a catechol-1,2-dioxygenase and deletion of NRRL3_04787 resulted in the inability to grow on catechol (Lubbers et al., 2021; Arentshorst and Ram, unpublished data). These results also show that the catabolism of gallic acid does not involve a pathway that includes protocatechuc acid or catechol as an intermediate.

The identification of the transcription factors controlling tannase expression also offers the possibility to optimize and increase the production of tannases. Tannases are biotechnologically interesting enzymes that are applied both in the clarification of wines, fruit juices and coffee-flavored drinks as well as in the production of gallic acid (Dhiman et al., 2018, Beniwal et al., 2013). As shown in this study, constitutive expression of certain tannases can be achieved by deleting the gene encoding the putative repressor protein tanX. Based on analogy with the GaaR/GaaX system, one can predict that overexpression of the gene encoding the TanR transcription factor will result in inducer-independent production of tannases, just as overexpression of gaaR led to inducer-independent expression of pectinases (Alazi et al., 2018).

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. RNA seq data regarding the parental strains (MA234.1) were previously deposited in the Short Read Archive under accession number SRP078415 (Niu et al., 2017). The RNA seq data regarding the ΔtanX mutant have been deposited in the Short Read Archive under accession number PRJNA713317. Metabolomics data have been deposited to the EMBL-EBI MetaboLights database (DOI: 10.1093/nar/gkz1019, PMID:31691833) with the identifier MTBLS2667.

**AUTHOR CONTRIBUTIONS**

MA: conceptualization, investigation, validation, data analysis, methodology, and visualization. MF: investigation, data analysis, methodology, investigation, and data analysis. M-CM: investigation and data analysis. IR: RNA sequencing and data analysis. TS and JD: investigation and data analysis. ED: conceptualization, data analysis, and manuscript draft writing. JP: conceptualization and data analysis. PP: conceptualization and manuscript draft writing. AT: conceptualization, data analysis, and funding. AR: conceptualization, data analysis, original draft writing, funding acquisition, and supervision. All authors contribute to the writing and agree to be accountable for the content of the work.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/ffunb.2021.681631/full#supplementary-material

**Supplementary Figure 1** | Southern blot analysis to verify tanX::hygR deletion in MA234.1 (A) Schematic representation of the tanX locus and the tanX::hygR locus in the deletion strain. DNA fragments expected to hybridize with the probe after digestion of genomic DNA with EcoRI are indicated. (B) Genomic DNA of putative ΔtanX::hygR transformants and the recipient strain MA234.1 was analyzed. MA586.1 was selected for further analysis.

**Supplementary Figure 2** | Diagnostic PCR to verify tanR::hygR deletion in MA234.1 (A) Schematic representation of the tanR::hygR locus in the deletion strain. Diagnostic PCR is performed using primer set 08275_P5f and hygR_5′ PCR and primer set hygR_3′ PCR and primer set 08275_P7f and 08275_P8r (3′ PCR). The locations where the primers anneal are indicated. (B) PCR reactions were performed with genomic DNA of putative tanR::hygR transformants as template and PCR products were analyzed using gel electrophoresis. MA514.1 was selected for further analysis.

**Supplementary Figure 3** | Diagnostic PCR to verify NRRL3_06002::AopyrG deletion in MA169.4 (A) Schematic representation of the NRRL3_06002 locus in the wild type (wt) strain and the NRRL3_06002::AopyrG locus in the deletion strain. Diagnostic PCR is performed using primer set 06002_P9f and 06002_P10r (wt 5′ PCR), primer set 06002_P7f and 06002_P8r (wt 3′ PCR), primer set 06002_P9f and AopyrGP16r (Δ 5′ PCR) and primer set AopyrGP17f and 06002_P8r (Δ 3′ PCR). The location where the primers anneal is indicated. (B) PCR reactions were performed with genomic DNA of a putative NRRL3_06002::AopyrG transformant and genomic DNA of wt strain N402 as template and PCR products were analyzed using gel electrophoresis. MA831.4 was used for further analysis.

**Supplementary Figure 4** | Diagnostic PCR to verify NRRL3_01651-56::AopyrG deletion in MA169.4. (A) Schematic representation of the NRRL3_01651-56 locus in the wild type (wt) strain and the NRRL3_01651-56::AopyrG locus in the deletion strain. Diagnostic PCR is performed using primer set 01651_P5f and 01651_P6r (wt 5′ PCR), primer set 01656_P7f and 01656_P8r (wt 3′ PCR), primer set 01651_P5f and AopyrGP16r (Δ 5′ PCR) and primer set AopyrGP17f and 01656_P8r (Δ 3′ PCR). The location where the primers anneal is indicated. (B) PCR reactions were performed with genomic DNA of twelve putative NRRL3_01651-56::AopyrG transformants and genomic DNA of wt strain N402 as template and PCR products were analyzed using gel electrophoresis. MA829.1 was used for further analysis.
Supplementary Figure 5 | Diagnostic PCR to verify NRRL3_02180-85::AopyrG deletion in MA169.4. (A) Schematic representation of the NRRL3_02180-85 locus in the wild type (wt) strain and the NRRL3_02180-85::AopyrG locus in the deletion strain. Diagnostic PCR is performed using primer set 02185_P5f and 02185_P6r (wt 5’ PCR), primer set 02185_P7f and 02185_P8r (wt 3’ PCR), primer set 02185_P5f and AopyrGP16r (A 5’ PCR) and primer set AopyrGP17f and 02185_P6r (A 3’ PCR). The location where the primers anneal is indicated. (B) PCR reactions were performed with genomic DNA of three putative NRRL3_02180-85::AopyrG transformants and genomic DNA of wt strain N402 as template and PCR products were analyzed using gel electrophoresis. MA852.1 was used for further analysis.

Supplementary Figure 6 | Diagnostic PCR to verify NRRL3_06147-50::AopyrG deletion in MA169.4. (A) Schematic representation of the NRRL3_06147-50 locus in the wild type (wt) strain and the NRRL3_06147-50::AopyrG locus in the deletion strain. Diagnostic PCR is performed using primer set 06147_P5f and 06147_P6r (wt 5’ PCR), primer set 06150_P7f and 06150_P8r (wt 3’ PCR), primer set 06147_P5f and AopyrGP16r (A 5’ PCR) and primer set AopyrGP17f and 06150_P6r (A 3’ PCR). The location where the primers anneal is indicated. (B) PCR reactions were performed with genomic DNA of four putative NRRL3_06147-50::AopyrG transformants and genomic DNA of wt strain N402 as template and PCR products were analyzed using gel electrophoresis. TS0.1 was used for further analysis.

Supplementary Figure 7 | Diagnostic PCR to verify NRRL3_08290-92::AopyrG deletion in MA169.4. (A) Schematic representation of the NRRL3_08290-92 locus in the wild type (wt) strain and the NRRL3_08290-92::AopyrG locus in the deletion strain. Diagnostic PCR is performed using primer set 08290_P5f and 08290_P6r (wt 5’ PCR), primer set 08292_P7f and 08292_P8r (wt 3’ PCR), primer set 08290_P5f and AopyrGP16r (A 5’ PCR) and primer set AopyrGP17f and 08292_P6r (A 3’ PCR). The location where the primers anneal is indicated. (B) PCR reactions were performed with genomic DNA of two putative NRRL3_08290-92::AopyrG transformants and genomic DNA of wt strain N402 as template and PCR products were analyzed using gel electrophoresis. MA857.2 was used for further analysis.

Supplementary Figure 8 | Diagnostic PCR to verify NRRL3_09054-57::AopyrG deletion in MA169.4. (A) Schematic representation of the NRRL3_09054-57 locus in the wild type (wt) strain and the NRRL3_09054-57::AopyrG locus in the deletion strain. Diagnostic PCR is performed using primer set 09054_P3f and 09054_P4r (wt 5’ PCR), primer set 09057_P3f and 09057_P4r (wt 3’ PCR), primer set 09054_P3f and AopyrGP16r (A 5’ PCR) and primer set AopyrGP17f and 09057_P4r (A 3’ PCR). The location where the primers anneal is indicated. (B) PCR reactions were performed with genomic DNA of two putative NRRL3_09054-57::AopyrG transformants and genomic DNA of wt strain N402 as template and PCR products were analyzed using gel electrophoresis. MA963.4 was used for further analysis.

Supplementary Figure 9 | Diagnostic PCR to verify NRRL3_10365-75::AopyrG deletion in MA169.4. (A) Schematic representation of the NRRL3_10365-75 locus in the wild type (wt) strain and the NRRL3_10365-75::AopyrG locus in the deletion strain. Diagnostic PCR is performed using primer set 10365_P5f and 10365_P6r (wt 5’ PCR), primer set 10375_P7f and 10375_P8r (wt 3’ PCR), primer set 10365_P5f and AopyrGP16r (A 5’ PCR) and primer set AopyrGP17f and 10375_P6r (A 3’ PCR). The location where the primers anneal is indicated. (B) PCR reactions were performed with genomic DNA of a putative NRRL3_10365-75::AopyrG transformant and genomic DNA of wt strain N402 as template and PCR products were analyzed using gel electrophoresis. MA960.2 was used for further analysis.

Supplementary Figure 10 | Diagnostic PCR to verify NRRL3_09659::AopyrG and NRRL3_04277::hygR deletion mutants. (A) Schematic representation of the NRRL3_09659::AopyrG locus in the wild type (wt) strain and the NRRL3_09659::AopyrG locus in the deletion strain. Diagnostic PCR is performed using primer set 09632_P5f and 09632_P6r (wt 5’ PCR), primer set 09632_P7f and 09632_P8r (wt 3’ PCR), primer set 09632_P5f and AopyrGP16r (A 5’ PCR) and primer set AopyrGP17f and 09632_P6r (A 3’ PCR). The location where the primers anneal is indicated. (B) PCR reactions were performed with genomic DNA of a putative NRRL3_09659::AopyrG transformant and genomic DNA of wt strain N402 as template and PCR products were analyzed using gel electrophoresis. TS0.1 was used for further analysis. (C) Schematic representation of the NRRL3_04277::hygR locus in the wild type (wt) strain and the NRRL3_04277::hygR locus in the deletion strain. Diagnostic PCR is performed using primer set 04277_P5f and 04277_P6r (wt 5’ PCR), primer set 04277_P7f and 04277_P8r (wt 3’ PCR), primer set 04277_P5f and hygRP16r (A 5’ PCR) and primer set hygRP2f and 04277_P8r (A 3’ PCR). The location where the primers anneal is indicated. (D) PCR reactions were performed with genomic DNA of a putative NRRL3_04277::hygR transformant and genomic DNA of wt strain N402 as template and PCR products were analyzed using gel electrophoresis. TS0.4 was used for further analysis.

Supplementary Figure 11 | Diagnostic PCR to verify deletion of 17 putative ring cleavage enzymes in MA234.1. (A) Schematic representation of a ring cleavage enzyme (RCE) locus in the wild type (wt) strain and the RCE locus in the deletion strain. The location where the primers anneal is indicated. (B) For each RCE deletion, a diagnostic PCR is performed using the primer sets shown in the table. Predicted lengths of PCR products for both the wt locus and the deletion locus are also shown. (C) PCR reactions were performed with genomic DNA of two putative transformants with a deletion of the 17 RCEs and genomic DNA of wt strain N402 as template and PCR products were analyzed using gel electrophoresis. MA962.5 was used for further analysis.

Supplementary Figure 12 | Growth analysis of A. niger reference strain (MA234.1), ΔtanR, ΔtanX, Δcse 17x, and ΔNRRL3_04659 on various carbon sources. Spores were point inoculated on minimal medium (MM) supplemented with the indicated carbon source (5 mM). Strains were grown for 5 days at 30°C before pictures were taken.

Supplementary Figure 13 | Growth analysis of A. niger reference strain (MA234.1), ΔtanR, ΔgacA strains on 2.5 mM fructose (left), 2.5 mM fructose and 25 mM gallic acid (middle), or 25 mM gallic acid (right). Spores were point inoculated on minimal medium (MM) supplemented with the indicated carbon source. Strains were grown for 3 days at 30°C before pictures were taken.

Supplementary Table 1 | Overview of the primers used in this study.

Supplementary Table 2 | TPM expression values.

Supplementary Table 3 | Gene clusters higher expressed in ΔtanX.

Supplementary Table 4 | Expression values of genes encoding ring cleavage enzymes.

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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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