Network Analysis Reveals Different Cellulose Degradation Strategies Across *Trichoderma harzianum* Strains Associated With XYR1 and CRE1

**Rafaela Rossi Rosolen**1,2, **Alexandre Hild Aono**1,2, **Déborah Aires Almeida**1,2, **Jaire Alves Ferreira Filho**1,2, **Maria Augusta Crivelente Horta**1 and **Anete Pereira De Souza**1,3*

1Center for Molecular Biology and Genetic Engineering (CBMEG), University of Campinas (UNICAMP), Campinas, Brazil, 2Graduate Program in Genetics and Molecular Biology, Institute of Biology, University of Campinas (UNICAMP), Campinas, Brazil, 3Department of Plant Biology, Institute of Biology, University of Campinas (UNICAMP), Campinas, Brazil

*Trichoderma harzianum*, whose gene expression is tightly controlled by the transcription factors (TFs) XYR1 and CRE1, is a potential candidate for hydrolytic enzyme production. Here, we performed a network analysis of *T. harzianum* IOC-3844 and *T. harzianum* CBMAI-0179 to explore how the regulation of these TFs varies between these strains. In addition, we explored the evolutionary relationships of XYR1 and CRE1 protein sequences among *Trichoderma* spp. The results of the *T. harzianum* strains were compared with those of *Trichoderma atroviride* CBMAI-0020, a mycoparasitic species. Although transcripts encoding carbohydrate-active enzymes (CAZymes), TFs, transporters, and proteins with unknown functions were coexpressed with *cre1* or *xyr1*, other proteins indirectly related to cellulose degradation were identified. The enriched GO terms describing the transcripts of these groups differed across all strains, and several metabolic pathways with high similarity between both regulators but strain-specific differences were identified. In addition, the CRE1 and XYR1 subnetworks presented different topology profiles in each strain, likely indicating differences in the influences of these regulators according to the fungi. The hubs of the *cre1* and *xyr1* groups included transcripts not yet characterized or described as being related to cellulose degradation. The first-neighbor analyses confirmed the results of the profile of the coexpressed transcripts in *cre1* and *xyr1*. The analyses of the shortest paths revealed that CAZymes upregulated under cellulose degradation conditions are most closely related to both regulators, and new targets between such signaling pathways were discovered.

**Abbreviations:** ABC, ATP-binding cassette; CAZymes, carbohydrate-active enzymes; CCR, carbon catabolite repression; CEs, carbohydrate esterases; CNT, concentrative nucleoside transporter; CRE1, carbon catabolite repressor 1; GHs, glycoside hydrolases; GO, Gene Ontology; GTs, glycosyltransferases; HRR, highest reciprocal rank; iTOL, Interactive Tree of Life; ITS, internal transcribed spacer; JTT, Jones-Taylor-Thornton; K2P, Kimura two-parameter; KEGG, Kyoto Encyclopedia of Genes and Genomes; KO, KEGG Orthology; MEGA, Molecular Evolutionary Genetics Analysis; MFS, major facilitator superfamily; ML, Maximum likelihood; PLs, polysaccharide lyases; Ta0020, *Trichoderma atroviride* CBMAI-0020; tef1, translational elongation factor 1; TFs, transcription factors; Th0179, *Trichoderma harzianum* CBMAI-0179; Th3844, *Trichoderma harzianum* IOC-3844; TOM, topological overlap matrix; TPM, transcripts per million; WGCNA, weighted correlation network analysis; XYR1, xylanase regulator 1.
Although the evaluated *T. harzianum* strains are phylogenetically close and their amino acid sequences related to Xyr1 and Cre1 are very similar, the set of transcripts related to *xyr1* and *cre1* differed, suggesting that each *T. harzianum* strain used a specific regulation strategy for cellulose degradation. More interestingly, our findings may suggest that Xyr1 and Cre1 indirectly regulate genes encoding proteins related to cellulose degradation in the evaluated *T. harzianum* strains. An improved understanding of the basic biology of fungi during the cellulose degradation process can contribute to the use of their enzymes in several biotechnological applications and pave the way for further studies on the differences across strains of the same species.

**Keywords:** Trichoderma harzianum, Trichoderma atroviride, cellulose, transcription factors, coexpression networks

### 1 INTRODUCTION

Lignocellulosic biomass is a complex recalcitrant structure that requires a consortium of carbohydrate-active enzymes (CAZymes) for its complete depolymerization. Due to their unique ability to secrete these proteins efficiently, filamentous fungi, such as *Trichoderma* spp. and *Aspergillus* spp., are widely explored for the industrial production of CAZymes (de Assis et al., 2015; Bischof et al., 2016). In the genus *Trichoderma*, *Trichoderma reesei* is the primary fungal industrial source of cellulases and hemicellulases (Martinez et al., 2008), while *Trichoderma harzianum* and *Trichoderma atroviride* have been widely explored for the industrial production of CAZymes (de Assis et al., 2015; Bischof et al., 2016). In the genus *Trichoderma*, *Trichoderma reesei* evolved by vertical transduction to achieve CCR (Horta et al., 2019; Han et al., 2020). Cre1 is the only conserved TF throughout the fungal kingdom, suggesting a conserved mechanism for CCR in fungi (Adnan et al., 2017). Interestingly, the effect of *cre1* deletion on cellulose-feeding microorganisms depends on the species (Portnoy et al., 2011; Mello-de-Sousa et al., 2014), whereas in *T. harzianum* P49P11, it can increase the expression levels of CAZyme genes and *xyr1* (Delabona et al., 2021).

Although *T. harzianum* strains have shown high cellulosytic activity (Horta et al., 2018; Li J. X. et al., 2020), most studies investigating this species have focused on biological control (Maruyama et al., 2020; Yan and Khan 2021). Thus, the regulatory mechanisms underlying hydrolytic enzyme production by these fungi are still poorly explored at the transcriptional level (Delabona et al., 2017; Delabona et al., 2020; Delabona et al., 2021). In addition, given the high degree of genetic variation observed within the genus *Trichoderma* (Kubicek et al., 2019) along with the complex speciation observed in the *T. harzianum* species (Druzhinina et al., 2010), it is necessary to explore how the main regulators Xyr1 and Cre1 behave across *T. harzianum* strains.

Previously, the transcriptomes of *T. harzianum* IOC-3844 (Th3844), *T. harzianum* CBMAI-0179 (Th0179), and *T. atroviride* CBMAI-0020 (Ta0020) were investigated under cellulose degradation conditions (Almeida et al., 2021). Different types of enzymatic profiles were reported, and both *T. harzianum* strains had higher cellulase activity than *T. atroviride*. Using this dataset, we aimed to investigate how the regulation of the TFs Cre1 and Xyr1 varies among *T. harzianum* strains. Based on the assumption that coexpressed genes tend to share similar expression patterns and that they could be coregulated by the same elements, we modeled a network of Th3844, Th0179, and Ta0020 using a weighted correlation network analysis (WGCNA) (Langfelder and Horvath 2008). The last strain, which is distantly related to *T. reesei* (Druzhinina et al., 2006) and represents a well-defined phylogenetic species (Dodd et al., 2003), was used to assess the differences across *Trichoderma* species. In addition, phylogenetic
analyses of XYR1 and CRE1 protein sequences were performed to clarify the evolutionary relationships of these regulators among the evaluated strains.

In this study, we identified and compared modules, hub genes, and metabolic pathways associated with CRE1 and XYR1 under cellulose degradation conditions. To deeply investigate their regulatory activities, we also performed first neighbor and shortest-path network analyses. Although the evaluated *T. harzianum* strains are phylogenetically close, by comparing their coexpressed transcript profiles, functional diversity was observed. This difference was accentuated when associating the results of Th0179 and Th3844 with those of Ta0020. Thus, we observed a specific transcriptional pattern related to CRE1 and XYR1 in each strain. Our study could contribute to improving our understanding of the regulation of cellulose degradation in *T. harzianum* and paves the way for further studies evaluating differences across strains within the same species. Addressing these questions by investigating the genetic regulatory mechanisms involved in cellulose degradation is important for enhancing both our basic understanding and biotechnological industrial applications of fungal abilities.

2 MATERIALS AND METHODS

2.1 Fungal Strains, Culture Conditions and Transcription Profiling

The species were obtained from the Brazilian Collection of Environment and Industry Microorganisms (CBMAI) located in the Chemical, Biological, and Agricultural Pluridisciplinary Research Center (CPQBA) of the University of Campinas (UNICAMP), Brazil. The identity of the *Trichoderma* isolates was authenticated by CBMAI based on phylogenetic studies of their internal transcribed spacer (ITS) region and translational elongation factor 1 (*tef1*) marker gene. The *T. harzianum* CBMAI-0179 (Th0179), *T. harzianum* IOC-3844 (Th3844), and *T. atroviride* CBMAI-0020 (Ta0020) strains were cultivated in simple carbon sources, including cellulose and glucose, to induce a clear genetic response. Details on the culture conditions and transcription profiling of the described experiments are provided in Supplementary Material S1, while the profile of enzymatic activities (cellulase, beta-glucosidase, and xylanase), the total protein content, and the constitution of the secreted proteome has been published previously (Horta et al., 2018; Almeida et al., 2021).

2.2 Phylogenetic Analyses

The ITS nucleotide sequences of *Trichoderma* spp. were retrieved from the NCBI database (https://www.ncbi.nlm.nih.gov/). Additionally, ITS nucleotide sequences amplified from the genomic DNA of Th3844, Th0179, and Ta0020 via PCR were kindly provided by the CBMAI and included in the phylogenetic analysis. The ITS region has been found to be among the markers with the highest probability of correctly identifying a very broad group of fungi (Schöch et al., 2012). *T. harzianum* T6776 was used as a reference genome to retrieve the CRE1 and XYR1 protein sequences belonging to *Trichoderma* spp. Additionally, the CRE1 and XYR1 sequences were obtained from the *T. harzianum* T6776 (Baroncelli et al., 2015) reference genome of Th3844 and Th0179 and the *T. atroviride* IMI206040 (Kubicke et al., 2011) genome of Ta0020 and used as references for the TF consensus sequences for RNA-Seq read mapping. These sequences were included in the phylogenetic analyses.

The multiple sequence alignment was performed using ClustalW (Thompson et al., 1994), and a phylogenetic tree was created using Molecular Evolutionary Genetics Analysis (MEGA) software v7.0 (Kumar et al., 2016). The maximum likelihood (ML) (Jones et al., 1992) method of inference was used based on (I) a Kimura two-parameter (K2P) model (Kimura 1980), (II) a Dayhoff model with frequencies (F’), and (III) a Jones-Taylor-Thornton (JTT) model for ITS, CRE1, and XYR1, respectively. We used 1,000 bootstrap replicates (Felsenstein 1985) in each analysis. The trees were visualized and edited using Interactive Tree of Life (ITOL) v6 (https://itol.embl.de/).

2.3 Weighted Gene Coexpression Network Analysis

The gene coexpression networks of Th3844, Th0179, and Ta0020 were modeled using transcripts per million (TPM) value data, which were validated through qPCR and described by Almeida et al. (2021), of three biological replicates with the R (R Core Team 2018) WGCNA package. Transcripts showing null values for most replicates under different experimental conditions were excluded. The network was assembled by calculating the Pearson’s correlation coefficient of each pair of genes. A soft power β was chosen for each network using pickSoftThreshold to fit the signed network to a scale-free topology. Then, an adjacency matrix in which the nodes correspond to transcripts and the edges correspond to the strength of their connection was obtained. To obtain a dissimilarity matrix, we built a topological overlap matrix (TOM) as implemented in the package.

To identify groups of transcripts densely connected in the gene coexpression network, we applied simple hierarchical clustering to the dissimilarity matrix. From the acquired dendrogram, the dynamicTreeCut package (Langfelder et al., 2008) was used to obtain the ideal number of groups and the respective categorization of the transcripts. According to the functional annotation performed by Almeida et al. (2021), groups containing the desired TFs XYR1 and CRE1 were identified and named the *xyr1* and *cre1* groups, respectively.

2.4 Functional Annotation of Transcripts in the *xyr1* and *cre1* Groups

We functionally annotated the transcripts in the *xyr1* and *cre1* groups. By conducting a Fisher’s exact test to extract the overrepresented terms (p-value < 0.05), all identified Gene Ontology (GO) (Ashburner et al., 2000) categories were used to identify enriched GO terms with the topGO package in R (Alexa and Rahnenführer 2021). To visualize the possible correlated enriched categories in the dataset caused by cellulose degradation, we created a treemap using the REVIGO.
tool (Supek et al., 2011). Then, the metabolic pathways related to the Kyoto Encyclopedia of Genes and Genomes (KEGG) (Kanehisa and Goto 2000) Orthology (KO) identified in T. reesei were selected due the high number of annotations correlated with this species in the KEGG database. To identify the pathways related to the enzymes identified as belonging to different groups, we used the Python 3 programming language (Sanner 1999) along with the BioPython library (Cock et al., 2009). The automatic annotation was manually revised using the UniProt databases (UniProt Consortium 2019).

2.5 Assessing the Groups’ Network Topologies
To provide a visual network characterization, we modeled additional gene coexpression networks of all strains using the highest reciprocal rank (HRR) approach (Mutwil et al., 2010). Using an R Pearson correlation coefficient threshold of 0.8, we assessed the 30 strongest edges, which were coded as ⅕ (top 10), 1/15 (top 20), and 1/30 (top 30). To visualize the behavior of the groups modeled by WGCNA in the HRR networks, we used Cytoscape software v3.7.0 (Shannon et al., 2003). Using the HRR groups modeled by WGCNA in the HRR networks, we used 15 (top 20), and 1/30 (top 30). To visualize the behavior of the transcripts, we calculated Pearson correlation matrices. Additionally, to represent the evolutionary relationships of CRE1 and XYR1 among Th3844, Th0179, and Ta0020, a phylogenetic analysis was performed while considering their amino acid sequences (Figure 1). The phylogenetic trees were modeled based on 26 and 21 protein sequences related to CRE1 and XYR1, respectively, in the Trichoderma genus, including those of our studied strains. Fusarium spp. was used as an outgroup. The NCBI accession number of the sequences used to model the phylogenetic trees is available in Supplementary Table S2.

The CRE1 phylogenetic tree indicated a close genetic proximity between Th3844 and Th0179 (with bootstrap support of 67%) (Figure 1A). These results were supported by the alignment of both protein sequences, which showed high conservation in the alignment of their amino acid sequences (with a percent identity of 99.76%) (Supplementary Figure S2). In contrast, the CRE1 protein sequence of Ta0020 shared closer affinity with two other T. atroviride strains (T23 and IMI206040), with bootstrap support of 99% (Figure 1A). Furthermore, the C2H2 domain is highly conserved in the Th3844, Th0179, and Ta0020 sequences of CRE1, with only one amino acid change from T. atroviride compared to both T. harzianum strains (Supplementary Figure S2).

In the XYR1 phylogenetic tree, Th3844, Th0179, and Ta0020 were closest to T. harzianum CBS 226.95 (with bootstrap support of 97%), T. harzianum T6776 (with bootstrap support of 51%), and T. atroviride IMI206040 (with bootstrap support of 88%), respectively (Figure 1B). Compared to the results of CRE1, the alignment of their amino acid sequences showed a lower percentage of identity, supporting the phylogenetic results observed (Supplementary Figure S2). In addition, the multiple alignments of the XYR1 protein sequence of Th3844, Th0179, and Ta0020 showed reasonable conservation of the Zn2Cys6 and fungal-specific TF domains (Supplementary Figure S2).

3.2 Weighted Gene Coexpression Network Analysis
Recently, the transcriptomes of Th3844, Th0179, and Ta0020 grown on crystalline cellulose and glucose after a time course of 96 h were investigated (Almeida et al., 2021). By using these transcriptome data and applying the described filters in the WGCNA package, we obtained (I) 11,050 transcripts (Th3844), (II) 11,105 transcripts (Th0179), and (III) 11,021 transcripts (Ta0020) (Supplementary Table S3). Based on these transcripts, we calculated Pearson’s correlation matrices. Then, different soft power β values were chosen (48 for Th3844, 8 for Th0179, and 27 for Ta0020) to obtain the scale-free topology, reaching fit indexes of (I) 0.8 for Th3844 (mean connectivity of 40), (II) 0.9 for Th0179 (mean connectivity of 842), and (III) 0.85 for Ta0020 (mean connectivity of 406). Then, the networks were partitioned into manageable groups to explore the putative coregulatory relationships. In total, we identified 87 groups in
Th3844, 75 groups in Th0179, and 100 groups in Ta0020 (Supplementary Table S4). Transcripts with expression patterns correlated with XYR1 and CRE1 were identified in the Th3844, Th0179, and Ta0020 coexpression networks and grouped (described in Supplementary Table S5). Among the strains, Ta0020 presented the highest number of transcripts coexpressed with cre1 and the lowest number of transcripts coexpressed with xyr1; the other strains showed the opposite profile.

3.3 Functional Characterization of the Transcripts in the xyr1 and cre1 Groups

We performed an enrichment analysis of the transcripts in each cre1 and xyr1 group of all evaluated strains using the GO categories (Supplementary Figures S3–S5, respectively). The cre1 group of Th3844, Th0179, and Ta0020 presented 50, 38, and 50 enriched GO terms, respectively, while in the xyr1 group, 34, 86, and 50 enriched GO terms were found in Th3844, Th0179, and Ta0020, respectively. Overall, the GO terms describing the transcripts of these groups were diverse across all strains, indicating a high degree of differences among the biological processes related to XYR1 and CRE1.

For example, the enrichment analyses of Ta0020 indicated that the carbohydrate metabolic process might be directly affected by CRE1 (high percentage of enriched GOs). Furthermore, GO terms related to fungal growth and nucleoside transmembrane transport were observed. In contrast, in Th0179 and Th3844, organic substance metabolic processes and regulation processes were pronounced, respectively. Interestingly, response to light stimulus was an enriched GO term in the cre1 group of Th3844, and fungal-type cell wall organization was an enriched GO term in Th0179. In the XRY1 group, Ta0020 also presented carbohydrate metabolic process as an enriched GO term. However, few genes corresponding to this term were observed, while terms related to the regulation of DNA transcription were pronounced. Interestingly, response to external stimulus and oxidation-reduction process were notable terms in Th0179 and Th3844, respectively. Furthermore, our results suggest that the regulation of biological processes was a common enriched term in all the evaluated strains, with some particularities across the T. harzianum strains, including regulation of cell population proliferation (Th0179) and mRNA metabolic process (Th3844).

The KO functional annotation of the transcripts encoding enzymes and unknown proteins with enzymatic activity in the cre1 and xyr1 groups was also investigated. In the 14 pathway classes observed in each TF, 13 were shared between the two regulators (Figure 2). Only two pathway classes, i.e., metabolism of other amino acids and lipid metabolism, were exclusive to CRE1 and XYR1, respectively. However, the number of identified enzymes in a given pathway class differed in each strain. This difference stresses the diversity of proteins with several functions, which could result in exclusive enzymatic performance in cellulose degradation according to the strain.

Next, we summarize some important aspects of the identified metabolic pathways. For example, Ta0020 showed a greater number of enriched pathway classes in the cre1 group, while both T. harzianum strains presented the opposite profile, with more enzymatic activity pathways enriched in the xyr1 group. Carbohydrate metabolism was a pathway class enriched in both T. harzianum strains in the xyr1 group, while Th0179 and Ta0020 presented such a profile in the cre1 group. In the xyr1 group, nucleotide metabolism and metabolism of cofactors and vitamins were enriched terms in Th0179, while xenobiotic degradation and
metabolism of terpenoids and polyketides were enriched terms exclusively in Ta0020. The pathway classes related to secondary metabolite compounds were also enriched in all evaluated strains in the cre1 and xyr1 groups, and Ta0020 presented the highest number of enzymes related to this term in the cre1 group.

In this study, network analyses were conducted to investigate how the molecular basis of XYR1 and CRE1 for cellulose-degrading enzyme production varies among T. harzianum strains. Thus, after analyzing the functional composition of the transcripts in the cre1 and xyr1 groups of all evaluated strains, particular attention was given to CAZymes, TFs, and transporters (Figure 3). We observed that in the cre1 group, Ta0020 showed the highest number of transcripts encoding CAZymes, TFs, and transporters, while in the xyr1 group, both T. harzianum strains presented a great number of coexpressed transcripts encoding TFs, followed by those encoding transporters. Furthermore, Th3844 showed a high number of CAZymes in the xyr1 group.

We identified the main CAZyme classes, including glycoside hydrolases (GHs), carbohydrate esterases (CEs), glycosyltransferases (GTs), and polysaccharide lyases (PLs) (Lombard et al., 2014), coexpressed with xyr1 and cre1 transcripts. To determine the similarities and differences across the strains, their CAZyme profiles were compared (Figure 4). In the xyr1 and cre1 groups, the GH family, which encompasses enzymes that hydrolyze glycosidic bonds, was identified in all the evaluated strains with different numbers of transcripts (Figures 4A,B). In the cre1 group, Ta0020 presented the highest number of classified transcripts of the GH family, followed by Th0179 and Th3844. In the xyr1 group, Th3844 presented a higher number of GHs than Th0179 and Ta0020. Furthermore, transcripts encoding CEs, which hydrolyze ester bonds, were associated with the cre1 transcript in Ta0020 and Th0179, whereas transcripts of the PL family, which cleave bonds in uronic acid-containing polysaccharide chains, were

**FIGURE 2** | KO functional classification of the transcripts identified in the cre1 and xyr1 groups. The sequences related to the enzymes coexpressed with cre1 (A) and xyr1 (B) were annotated according to the main KO functions of Th3844, Th0179, and Ta0020. For all evaluated strains, 14 pathway classes were observed in each TF, and 13 classes were shared between the two regulators. However, the number of identified enzymes in a given pathway class differed in each strain. Each strain is represented by a different color. Ta0020: T. atroviride CBMAI-0020; Th0179: T. harzianum CBMAI-0179; Th3844: T. harzianum IOC-3844; KO: Kyoto Encyclopedia of Genes and Genomes Orthology.
coexpressed with the xyr1 transcript in only Th3844. In addition, transcripts of the GT family, which synthesizes glycosidic bonds from phosphate-activated sugar donors, were present in the cre1 group of Th0179 and the xyr1 groups of Th3844 and Ta0020. We also investigated the quantification of each CAZyme family in the cre1 and xyr1 groups of all evaluated strains (Figures 4C,D). Overall, Ta0020 presented a high number of GHs belonging to family 18, coexpressed with the cre1 transcript, while Th0179 exhibited a significant amount of GT90 (Figure 4C). Different CAZyme profiles were observed in the xyr1 groups, and Th3844 showed the highest diversity of CAZyme families (Figure 4D).

Several positive and negative regulators are involved in the expression of CAZyme-coding genes and other proteins required for lignocellulose breakdown. Thus, we sought to identify transcripts encoding TFs coexpressed in the cre1 and xyr1 groups (Figure 5). In both groups, transcripts encoding Zn2Cys6 type TFs were identified in all the evaluated strains with different numbers of transcripts. In the cre1 group of Ta0020, the number was higher than that observed in Th0179 and Th3844 (Figure 5A), while in the xyr1 group, we observed the opposite profile (Figure 5B). Most Zn2Cys6 proteins also contain a fungal-specific TF domain, which was coexpressed with both cre1 and xyr1 transcripts in Th3844 and Ta0020. In this last strain, a C6 zinc finger domain TF was coexpressed with xyr1. Transcripts encoding C2H2 type TFs were identified in Ta0020 and T. harzianum strains in the cre1 and xyr1 groups, respectively. Furthermore, in the xyr1 group, we found a transcript encoding the SteA regulator, which is involved in the regulation of fungal development and pathogenicity (Hoi and Dumas 2010), in Th0179.

Transporters are also important players in the degradation of plant biomass. Here, transcripts encoding transport proteins coexpressed with cre1 and xyr1 were selected (Figure 5). Among them, transcripts encoding major facilitator superfamily (MFS) transport proteins were coexpressed with cre1 in Th0179 and Ta0020 (Figure 5C) and xyr1 in Th0179 and Th3844 (Figure 5D). Amino acid transporters were present in the xyr1 groups of all strains and the cre1 group only of Th0179. Transcripts encoding the ATP-binding cassette (ABC) were coexpressed with xyr1 only in Th3844. Th0179 showed transcripts encoding several types of transporters coexpressed with xyr1, e.g., vesicle transporter SEC22, cation transporting ATPase, and calcium-dependent mitochondrial carrier, while Ta0020 coexpressed with cre1, e.g., UDP-galactose transporter and concentrative nucleoside

FIGURE 3 | Comparisons of TFs, transporters, and CAZymes among strains and groups. Number of transcripts encoding TFs, transporters, and CAZymes in Ta0020, Th0179, and Th3844 distributed among the cre1 (A) and xyr1 (B) groups. The number of transcripts encoding these proteins associated with cre1 was more accentuated in Ta0020 than those observed for the other strains, while in the xyr1 group, both T. harzianum strains presented a great number of coexpressed transcripts encoding TFs, followed by those encoding transporters. Th3844 also showed a high number of CAZymes in the xyr1 group. TFs: transcription factors; CAZymes: carbohydrate-active enzymes. Ta0020: T. atroviride CBMAI-0020; Th0179: T. harzianum CBMAI-0179; Th3844: T. harzianum IOC-3844.
transporter (CNT). Transcripts encoding ion transporters were present in both the cre1 and xyr1 groups of Ta0020 and Th0179. In the cre1 group, such transcripts included zinc and magnesium transporter proteins in Ta0020 and potassium transporter proteins in Th0179. In the xyr1 group, such transcripts include Cd²⁺Zn²⁺ transporters in Ta0020 and Ca²⁺ transporters in Th0179. Th0179 and Th3844 also present a G protein coexpressed with the cre1 transcript.

To better understand the regulation mediated by the studied TFs in the evaluated T. harzianum strains, we also investigated other classes of proteins coexpressed with the cre1 and xyr1 transcripts. For example, phosphatases, kinases and TFs are key
components in cellular signaling networks. Transcripts encoding kinase proteins were coexpressed with \textit{cre1} transcripts in Ta0020 and Th3844 and \textit{xyr1} transcripts in Ta0020 and Th0179. In addition, all strains showed transcripts encoding phosphatases coexpressed with \textit{cre1}, and in Th0179 and Th3844, such transcripts were coexpressed with \textit{xyr1}. In addition, transcripts encoding cytochrome P450 genes, which constitute an important group of enzymes involved in xenobiotic degradation and metabolism, were found in the \textit{cre1} groups of all the evaluated strains.

Furthermore, different quantities of differentially expressed transcripts were found in Ta0020 (78) and Th0179 (6) (described in Supplementary Tables S6, S7). In Ta0020, most upregulated transcripts under cellulose growth (70) were associated with the \textit{cre1} group, including \textit{cre1}, whereas \textit{xyr1} demonstrated a potentially significant expression level under glucose growth. In Th3844, \textit{cre1} significantly modulated the expression of the two carbon sources, with a high expression level under glucose growth. Even with the reduced differentiation observed (−1.13), this phenomenon was considered in other studies (Antonieto et al., 2014; Castro et al., 2016). In Th0179, \textit{xyr1} and \textit{cre1} revealed a low modulation in transcript expression between growth under cellulose or glucose. However, \textit{xyr1} had a higher transcript expression level under glucose than \textit{cre1}, which had a higher transcript expression level under cellulose growth conditions. In Th3844, \textit{xyr1} exhibited similar results.

3.4 Assessing the Groups’ Network Topologies
To identify the main aspects of the network topologies, we applied the HRR methodology to the transcriptome data,
which were also used to model the networks using the WGCNA methodology (Figures 6A–C).

In the CRE1 and XYR1 subnetworks, different connection profiles were observed in each strain. While the transcripts in the crel groups showed less ranked connections in the WGCNA groups of the T. harzianum strains Figures 6D,E), Ta0020 showed the opposite profile. In this strain, the transcripts in the crel group were more strongly connected (Figure 6F). In contrast, in the xyrl groups, the transcripts were more densely connected in the T. harzianum strains (Figures 6G,H), while in Ta0020, the transcripts were separated (Figure 6I).

Transcripts at the top of the degree distribution, which are defined here as hubs, are topologically important to the network structure and are often functionally relevant (Luscombe et al., 2004). Thus, the hub nodes were sought in the crel and xyrl groups. In the crel groups, transcripts encoding a SET-domain protein, an ATP synthase, and a transcript not yet annotated were encoded hypothetical proteins, while in the xyrl groups, the transcripts were distributed among 24 groups (Supplementary Table S8).

In Ta0020, we found that such an uncharacterized transcript had a degree value equal to 52, which differs from the T. harzianum strains in which the hub nodes had degree values of 4 (Th3844) and 9 (Th0179) (Supplementary Table S8). In the xyrl groups, transcripts encoding an ATP-dependent RNA helicase, a cytokinesis sepA, and hypothetical proteins were found as hub nodes of Th3844, Th0179, and Ta0020, respectively. In Ta0020, we found that such an uncharacterized transcript had a degree value equal to 52, which differs from the T. harzianum strains in which the hub nodes had degree values of 4 (Th3844) and 9 (Th0179) (Supplementary Table S8). In the xyrl groups, transcripts encoding an ATP-dependent RNA helicase, a cytokinesis sepA, and hypothetical proteins were found as hub nodes of Th3844, Th0179, and Ta0020, respectively (Supplementary Table S8).

Here, we show that the hub nodes in the crel group of Ta0020 had a degree value of 1 and all encoded hypothetical proteins, while in the T. harzianum strains, the hub nodes had degree values of 13 (Th3844) and 7 (Th0179). Among the other transcripts with a degree value lower than that of those at the top of the degree distribution, in the xyrl group, GH76 and Zn2Cys6-type TFs (degree value of 6) were identified as candidate hub genes of Th0179, while regulator-nonsense transcripts (degree value of 9) and Zn2Cys6-type TFs (degree value of 8) were identified as candidate hub genes of Th3844. In the crel group, the proteasome component PRE3 (degree value of 2) in Th3844 was found as a hub node, and replication factor C and phosphoglucomutase (degree value of 5) were found in Th0179. In addition, in Ta0020, we identified calcium calmodulin-dependent kinase (degree value of 9) and GH93 (degree value of 8) as hub nodes.

In all evaluated strains, as a reduced number of differentially expressed transcripts was found in the crel and xyrl groups, we expanded our investigation of the studied groups by selecting the first neighbors of XYR1 and CRE1 in global HRR networks. Different quantities of crel neighbors were found in Th3844 (59), Th0179 (21), and Ta0020 (25), and different quantities of xyrl were found in Th3844 (46), Th0179 (59), and Ta0020 (70) (described in Supplementary Table S9). Regarding the crel neighbors, these transcripts were distributed among 24 groups in Th3844, 15 groups in Th0179, and 16 groups in Ta0020. In contrast, considering the xyrl neighbors, such transcripts were distributed among 19 groups in Th3844, 32 groups in Th0179, and 42 groups in Ta0020. The direct coexpressed neighbors of xyrl included downregulated transcripts in Th3844 (4), Th0179 (5), and Ta0020 (22) and upregulated transcripts in Th3844 (1), Th0179 (1), and Ta0020 (1). Interestingly, only Ta0020 showed differentially expressed transcripts as the first neighbor of the crel transcript (4 downregulated and 4 upregulated). Overall, these differentially expressed transcripts included transporters, CAZymes, kinases, and other proteins related to the regulation process.

For simplification, only a few neighboring transcripts of xyrl and crel in Th3844, Th0179, and Ta0020 are shown in Table 1. In Th3844, TFs, transporters, a cytochrome, and a phosphatase were found to be first neighbors of the crel transcript, while TFs, transporters, and CAZymes were found to be the first neighbors of the xyrl transcript. In Th0179, several types of proteins, including a kinase, were found to be first neighbors of crel, while transporters, CAZymes, and a TF were found to be first neighbors of the xyrl transcript. In Ta0020, several types...
of proteins, including kinases, were found to be first neighbors of cre1, while transporters, CAZymes, TFs, and a DNA ligase were found to be first neighbors of the xyr1 transcript. Hypothetical proteins were also found to be neighbors of the cre1 and xyr1 transcripts of all strains, indicating that both regulators might have a regulatory influence on uncharacterized proteins.

We also applied a shortest-path network analysis to identify possible transcripts outside of the xyr1 and cre1 groups that may be influenced by CRE1 and XRY1 (described in Supplementary Table S10). As the shortest path, we considered the minimal number of edges that need to be traversed from a node to reach another node (Koutrouli et al., 2020). As both regulators act on the gene expression of hydrolytic enzymes related to plant biomass degradation, we sought to determine the number of shortest paths from XYR1 and CRE1 to the selected CAZymes. In all evaluated strains and both TFs, we found a similar quantity of shortest paths (Figure 7A; Supplementary Figures S6, S7). However, some aspects should be noted. For example, Th0179 presented transcripts encoding CAZymes from the GH1, GH5, and GH6 families, which are more distantly related to CRE1, while Ta0020 showed only a CAZyme from the GH6 family satisfying this requirement. In Th3844, we found a transcript encoding a CAZyme from GH3 strongly related to XYR1, and CAZymes with the CBM1 domain were also closely connected with such a regulator. In Th0179, a CAZyme from GH55 exhibited a close relationship with XYR1, while in Ta0020, CAZymes from the GH7, GH72, and GH1 families and the CBM1 domain also presented such a profile. In addition, given a determined shortest path, we also considered the number of possibilities that may have occurred (Figure 7B; Supplementary Figures S6, S7). Across all strains and in both TFs, significant differences were observed in the number of chances. For instance, a significant number of possible minimum paths between CRE1 and a CAZyme from the GH6 family were observed in Ta0020, likely suggesting a strong influence of this repressor regulator on enzyme activity. Similarly, CAZymes from the AA9 and GH16 families appeared to be affected by XYR1. In Th0179, CAZymes from GH10, GH6, and GH1 presented a higher number of possible shortest paths related to CRE1, potentially indicating an impact of this TF on these enzyme activities.

We also performed a shortest-path analysis to identify transitive transcripts between cre1 or xyr1 and the selected CAZymes, which allowed us to discover new transcripts that may be involved in the same biological process. For simplification, we only chose a few shortest paths to explore such transitive transcripts. In Th3844, NADH-ubiquinone oxidoreductase (THAR02_08259), which is the largest multiprotein complex of the mitochondrial respiratory chain (Whitehouse et al., 2019), was found between GH45 with a CBM1 domain (THAR02_02979) and cre1. We also found another transcript involved in ATP synthesis, NAD(P) transhydrogenase beta subunit (THAR02_08260), between GH1 (THAR02_05432) and cre1. Furthermore, a transcript encoding a component of the endoplasmic reticulum quality control system called ER-associated degradation (THAR02_08220) (Phillips et al., 2020) was found between GH16 (THAR02_03302) and cre1. Interestingly, in Ta0020, a phosphotyrosine phosphatase was found in the shortest path between GH1 (TRIATDRAFT_135426), which is a homolog of GH1 (THAR02_05432) in T. harzianum strains, and cre1. This phosphatase is involved in protein phosphorylation, which is a
key posttranslational modification critical for the control of many cellular functions (Nasa and Kettenbach 2018). In Th3844, we found a transcript encoding a mitochondrial outer membrane porin (THAR02_07078) between CBM1 (THAR02_02133) and xyr1. These protein transporters transport small molecules and play significant roles in diverse cellular processes, including the regulation of mitochondrial ATP and calcium flux (Grevel and Becker 2020). In Ta0020, methyltransferase domain-containing (TRIATDRAFT_292180), which is important for the regulation of chromatin and gene expression (Kouzarides 2007), was found between GH1 (TRIATDRAFT_150220) and xyr1.

4 DISCUSSION

Although the importance of XYR1 and CRE1 in the expression of CAZyme-encoding genes and other proteins required for lignocellulose degradation is evident, the transcriptional regulation mediated by both proteins in T. harzianum strains remains poorly explored (Delabona et al., 2017; Delabona et al., 2021). A previous study demonstrated that some genes encoding regulatory proteins, such as xyr1, evolved by vertical gene transfer in Trichoderma spp. (Druzhinina et al., 2018). Here, we inferred the evolutionary relationships of XYR1 and CRE1 in the evaluated strains. In both regulatory proteins, a great genetic distance was observed between the amino acid sequences of Ta0020 and those of Th3844 and Th0179. These findings supported the genetic tree of the evaluated species in which Ta0020 was grouped with other T. atroviride strains. However, among the T. harzianum strains, we noted that the CRE1 amino acid sequences were more highly conserved than those of XYR1. These results are consistent with previous studies reporting that while the function of CRE1 is conserved throughout the fungal kingdom (Adnan et al., 2017), the role of XYR1 greatly differs across ascomycete fungi (Klaubauf et al., 2014).

Recently, different types of enzymatic profiles across Trichoderma species were reported, and Th3844 and Th0179, which have hydrolytic potential, have higher cellulase activity during growth on cellulose than Ta0020 (Almeida et al., 2021). Because such diversity in the enzyme response can be affected by the specific functionalities of regulatory proteins and evolutionary divergence between XYR1 and CRE1 (Klaubauf et al., 2014; Benocci et al., 2017), we aimed to investigate how both TFs could affect the activities of the transcripts in response to cellulose degradation in the T. harzianum strains. Because transcripts that share the same function or are involved in the same regulatory pathway tend to present similar expression profiles and, hence, form modules (Wolfe et al., 2005), networks were modeled for Th3844, Th0179, and Ta0020, and the last strain was used to assess the differences across Trichoderma species. In fungi, such methodology has been successfully used to provide insight into the regulatory mechanisms of hydrolysis (Borin et al., 2018; Arntzen et al., 2020; Li C. X. et al., 2020).

In recent years, network methodologies have been employed for the discovery of new targets associated with biological processes (Cortijo et al., 2020; Liu et al., 2021; Zhang et al., 2021). Because such approaches might provide mechanisms for inferring new gene relationships and functions, experimental validation by the construction of mutants is usually a way of corroborating these insights. In this context, the construction of mutants can show the influence of a unique missing gene on genome regulation, helping to corroborate the hypotheses raised in the present study. However, due to the inefficient homologous recombination machinery of filamentous fungi, once a fungus reproduces asexually, a low frequency of correct genomic integration has been observed for Trichoderma spp. (Zeilinger 2004; Liu et al., 2017; Vieira et al., 2018). These challenges can be overcome by using the CRISPR/Cas9 system, which represents a good alternative in Trichoderma spp. (Fonseca et al., 2020; Vieira et al., 2021) and may enable the future establishment of efficient protocols for each strain described here. Thus, by predicting the profiles of the transcripts coexpressed with XYR1 and CRE1, the results can be used to engineer mutant strains with the objective to investigate the influence of the regulation of such TFs in vitro.

We observed that each network had a different configuration with distinct module profiles of XYR1 and CRE1, providing insight into how these TFs might act in specific ways in different Trichoderma species. These functional differences in the xyr1 and cre1 groups can be attributed to previously reported differences in the regulatory mechanisms of hydrolysis (Almeida et al., 2021). By analyzing both groups, we observed that more transcripts were coexpressed with XYR1 in the T. harzianum strains than T. atroviride, which had more transcripts coexpressed with CRE1. However, although the phylogenetic similarity between the evaluated T. harzianum strains shows that they share a largely common genetic background, their profiles of transcripts coexpressed with xyr1 and cre1 differed.

4.1 Insight Into the Genetic Impacts of XYR1 and CRE1

To obtain insight into the functional profile of the cre1 and xyr1 groups, GO enrichment analyses of transcripts from these groups of all evaluated strains were performed. Here, response to external stimulus was a notable GO term in Th0179 in the xyr1 group, suggesting that the transcripts of these group likely respond to external environmental conditions, such as carbon sources. Furthermore, interconnections between nutrient and light signaling pathways have been reported in filamentous fungi, such as N. crassa and T. reesei, with substantial regulation by photoreceptors (Schmoll 2018). More interestingly, the influence of light on CRE1 functions has been reported (Monroy et al., 2017), supporting our findings showing that the response to a light stimulus was an enriched GO term in the cre1 group of Th3844. In contrast, the enrichment analysis of Ta0020 indicates that the activity of CRE1 may be stronger in such fungi, directly repressing genes related to plant cell wall-degrading enzymes, which was not observed in the T. harzianum strains. In these strains, organic substance metabolic processes (Th3844) and
regulation processes (Th0179) were enriched GO terms. In addition, fungal-type cell wall organization was an enriched term of Th0179 in the cre1 group.

By investigating the KO functional annotation of the cre1 and xyr1 groups, we suggest that both regulators act on the same enzymatic pathways, which was expected due to antagonism in their function, i.e., while CRE1 is the main repressor of genes encoding proteins related to lignocellulose degradation, XRY1 is the main activator of such genes. However, each triggers a specific metabolic pathway according to the strain; therefore, some aspects are noteworthy. For example, it has been reported that the Trichoderma species display several mechanisms during their antagonistic action against plant pathogens, including the production of secondary metabolites (Malmierca et al., 2015). According to our findings, due to the highest number of pathways related to secondary metabolites, including the metabolism of terpenoids and polyketides (Mukherjee et al., 2008), we suggest that CRE1 and XYR1 participate in the regulation of secondary metabolism compounds in T. atroviride. Furthermore, XYR1 in T. harzianum strains appeared to be deeply connected with enzyme pathways related to the metabolism of carbohydrates, especially in the Th3844 strain.

Through a network analysis, we identified transcripts encoding CAZymes coexpressed with xyr1 and cre1 in all evaluated strains. We identified CAZyme families responsible for cellulose degradation (e.g., GH12); hemicellulose degradation (e.g., GH16, GH17, CE1, and CE5); pectin degradation (e.g., GH28 and GH93); and other CAZymes with multiple activities or minor activities on lignocellulosic substrates, such as GH2, GH5, GH43, GH95, GH30, and GH39 (de Vries et al., 2017; Kameshwar et al., 2019). Although Ta0020 presented the highest number of GHs coexpressed with CRE1, it was found to be less efficient than the other Trichoderma strains in degrading plant biomass (Almeida et al., 2021). In the GH class, enzymes belonging to the GH18 family, which is mainly represented by chitinase-like proteins, are directly related to fungal cell wall degradation in mycoparasite species of Trichoderma (Gruber and Seidl-Seiboth 2012). Four GH18 enzymes were coexpressed with CRE1 in T. atroviride, which is widely used as a biocontrol agent in agriculture. However, in Ta0020, enzymes from the GH75 family of chitosanases were coexpressed with xyr1. Therefore, the degradation of chitosan, which is a partially decacylated derivative of chitin (Hahn et al., 2020), is a relevant aspect of mycoparasitism that may be influenced by the activity of XYR1. Recently, transcripts encoding GHs were found to be coexpressed with some TFs involved in biomass degradation, e.g., XYR1 (Borin et al., 2018). In the present study, the transcripts encoding hydrolytic enzymes could be sorted into other groups formed in the networks mainly because their expression patterns differed from those of the xyr1 and cre1 transcripts.

Several fungal TFs have been described to be directly involved in the regulation of plant biomass utilization (Benocci et al., 2017). Most TFs belong to the zinc cluster family, including Zn2Cys6 and C2H2-type TFs, which are characterized by the presence of zinc finger(s) in their binding domains. Most positive regulators appear to belong to the Zn2Cys6 class, while repressors belong to the C2H2 class (Benocci et al., 2017). Here, we found the highest number of transcripts encoding the Zn2Cys6 and C2H2 classes in the xyr1 and cre1 groups of Th0179 and Ta0020, respectively. Both T. harzianum strains had a similar profile of transcripts coexpressed with CRE1 and XYR1. However, SteA, a C2H2-type TF, was coexpressed with the xyr1 transcript of Th0179. This regulator has been described as an important player in fungal environmental adaptation in response to nutrient deprivation, the production of extracellular proteins involved in the degradation of complex substrates (Hoi and Dumas 2010), and the mediation of the regulatory role of mitogen-activated protein kinase (MAPK) during mycoparasitic responses (Gruber and Zeilinger 2014). In Aspergillus nidulans, SteA is required for sexual development (Vallim et al., 2000).

During plant biomass degradation, fungi secrete extracellular enzymes to decompose polysaccharides into small molecules, which are then imported into cells through transporters (Sloothaak et al., 2016). One of the most relevant sugar transporter families in filamentous fungi is the MFS family (Zhang et al., 2013). Here, we found the highest number of transcripts encoding MFS coexpressed with cre1 and xyr1 in Ta0020 and Th3844, respectively. Among the T. harzianum strains, compared with Th3844, Th0179 showed transcripts encoding several types of transporter proteins, such as transporter proteins of calcium ions (THAR02_04202, THAR02_03989, and THAR02_01350). It has already been reported that metal ions, such as Ca2+, have a positive effect on the mycelial growth of T. reesei and cellulase production (Chen et al., 2016). This molecular signaling mechanism is mediated by cations transporting ATPase and calcium-dependent mitochondrial carriers, which are both components of Ca2+/calmodulin signal transduction, including the TF Crz1 (Chen et al., 2016; Martins-Santana et al., 2020). Therefore, investigating the role of proteins related to calcium transporters in the induction of genes responsive to lignocellulose degradation in Th0179 cells is important since these proteins could transport cations that activate gene expression. Furthermore, transcripts encoding G proteins were coexpressed with cre1 in Th0179 and Ta0020. Heterotrimeric G proteins have been well studied in several Trichoderma species. In saprophytic species, such proteins are involved in the nutrient signaling pathway in connection with a light response, triggering the posttranscriptional regulation of cellulase expression (Hinterdobler et al., 2021); in mycoparasitic species, G protein-coupled receptors are involved in the regulation of processes related to mycoparasitism (Zeilinger and Atanasova 2020).

Although CAZymes, TFs, and transporters play an important role in cellulose degradation, these types of proteins represented only a small percent of the transcripts coexpressed with XYR1 and CRE1 compared to the other protein classes as follows: in the cre1 group, (I) 10.4% (Th0179), (II) 5.7% (Th3844), and (III) 17.9% (Ta0020); and in the xyr1 group, (IV) 11.8% (Th0179), (V) 11.9% (Th3844), and (VI) 16% (Ta0020). Since the regulation of genes involved in biomass breakdown is a complex process that involves several signaling pathways, we expected to find proteins with a great range of functions coexpressed with xyr1 and cre1. For example, various kinases and phosphatases were coexpressed with xyr1 and cre1 in all
evaluated strains. It has been reported that cellulase gene expression can be regulated by the dynamics of protein phosphorylation and dephosphorylation, which involve protein kinases and phosphatases, respectively (Schmoll et al., 2016). Furthermore, in filamentous fungi, phosphorylation is a prerequisite for CRE1 activity (Cziferszky et al., 2002; Han et al., 2020; de Assis et al., 2021). Therefore, in T. harzianum, it is important to elucidate the role of kinases and phosphatases in the regulation of CRE1 function. In all evaluated strains, cytochrome P450 coding genes represented another class of proteins coexpressed in the cre1 group. It has been reported that these enzymes are important for cells to perform a wide variety of functions, including primary and secondary metabolism, xenobiotic degradation, and cellular defense against plant pathogenic fungi (Siewers et al., 2005; Fan et al., 2013; Chadha et al., 2018). To expand previous findings concerning Trichoderma spp. (Chadha et al., 2018), their similar expression pattern with the crel transcript makes them important candidates for an extensive investigation in the cellulose degradation context.

We also investigated the expression profiles of the transcripts in the crel and xyr1 groups, including xyr1 and crel, under cellulose and glucose growth. Castro et al. (2016) reported that in T. reesei, the expression level of xyr1 was minimal in the presence of glucose; this phenomenon was not observed in Ta0020 in the present study. In addition, in Ta0020, crel was upregulated in the presence of cellulose, which was not expected due to the role of CRE1 in the repression of cellulolytic and hemicellulolytic enzymes when an easily metabolizable sugar, i.e., glucose, is available in the environment (Benocci et al., 2017). In contrast, in Th3844, the crel expression level in the presence of glucose was higher than that in the presence of cellulose. The gene coexpression networks were modeled based on the transcript expression level under two sets of conditions (cellulose and glucose). Considering that the expression of the crel transcript in Ta0020 was upregulated under cellulose growth, upregulated transcripts were expected and found in the crel group. Recently, Almeida et al. (Almeida et al., 2021) reported that Ta0020 presented the highest number of differentially expressed transcripts under cellulose growth conditions relative to glucose, followed by Th3844 and Th0179. Here, the quantity of upregulated transcripts was low in the crel and xyr1 groups of the T. harzianum strains. These results may indicate that both TFs have a basal expression level at 96 h of fermentation; thus, the transcripts grouped with such regulatory proteins presented the same expression pattern and, mostly, were not differentially expressed transcripts.

4.2 Examining the Network’s Topology

To extract additional information from the networks, we characterized the network topologies of the modules identified separately. We might infer that many transcripts were under the influence of the repressor CRE1 in Ta0020, while in the T. harzianum strains, the transcripts seem to have been affected by the activator XYR1. Therefore, the described profiles may indicate that in T. harzianum, the transcripts, including those encoding XYR1, act together to perform a determined biological function that is favorable to the expression of genes related to cellulose degradation. In contrast, in T. atroviride, the transcripts in the crel group appeared to act on the same biological process, which may be related to the repression of genes encoding hydrolytic enzymes and other proteins required for cellulose degradation.

Hub transcripts were identified in the crel and xyr1 groups, and new targets were discovered. In the crel group, a transcript with a SET-domain coding gene was found as a hub node of Th3844. Such SET-domain proteins participate in chromatin modifications by methylation specific lysines on the histone tails (Kouzarides 2007). In filamentous fungi, the epigenetic regulation of holocellulase gene expression has already been reported (Zeilinger et al., 2003), and CRE1 plays an important role in nucleosome positioning (Ries et al., 2014). Interestingly, ATP synthase, which synthesizes ATP from ADP and inorganic phosphate on mitochondria (Burger et al., 2003), was found to be a hub node of Th0179. Thus, we might infer that a great number of transcripts were coexpressed with a hub node related to mitochondrial ATP production, which is the main energy source for intracellular metabolic pathways (Neupane et al., 2019). In the xyr1 group, a transcript encoding an ATP-dependent RNA helicase was found as a hub node of Th3844. Such enzymes catalyze the ATP-dependent separation of double-stranded RNA and participate in nearly all aspects of RNA metabolism (Jankowsky 2011). Additionally, the sepA transcript was identified as a hub node of Th0179. In Aspergillus nidulans, the sepA gene encodes a member of the FH1/2 protein family, which is involved in cytokinesis and the maintenance of cellular polarity, i.e., related to the cell division process (Harris et al., 1997). Overall, our findings suggest that the hub genes were not necessarily the most effective genes related to lignocellulose deconstruction, confirming the indirect action of XYR1 and CRE1 in T. harzianum on regulatory hydrolysis mechanisms. In Ta0020, transcripts encoding hypothetical proteins were identified as potential hub nodes in both groups, providing new targets for further studies evaluating their functions in fungal physiology.

We also investigated the first neighbors of the crel and xyr1 transcripts in the modeled global HRR networks of all evaluated strains. While Ta0020 showed differentially expressed transcripts as the first neighbor of the crel transcript, interestingly, both T. harzianum strains presented the opposite profile in which only the first neighbors of the xyr1 transcript were differentially expressed. In addition, the evaluated strains showed different numbers of neighbors of the crel and xyr1 transcripts, which were distributed among several groups. Such a divergent profile may indicate that each TF affects a set of transcripts in a specific way that varies among the strains. Overall, we identified CAZymes, TFs, and transporters as first neighbors of crel and xyr1, confirming the results obtained in this study using the WGCNA approach.

Another property of a network’s topology is the shortest paths connecting two transcripts (Pavlopoulos et al., 2011). Here, the significant shortest paths between both studied TFs, i.e., CRE1 and XYR1, and the CAZymes with a higher-level expression under cellulose growth conditions were investigated in all strains. The average shortest pathway distance between the transcripts...
In conclusion, biological networks represent a powerful approach to accelerate the elucidation of the molecular mechanisms underlying important biological processes. Our findings suggest that the set of transcripts related to XYR1 and CRE1 varies among the studied T. harzianum strains, suggesting regulatory differences in enzymatic hydrolysis. These findings corroborate previous studies in which differences in biomass degradation and enzyme production between strains of the same species have been reported (de Vries et al., 2017; Thanh et al., 2019; Tolgo et al., 2021). Furthermore, such transcripts were not limited to CAZymes and other proteins related to biomass degradation. Thus, we suggest that both TFs play a role in the undirected regulation of gene encoding proteins related to cellulose degradation, and multiple pathways related to gene regulation, protein expression, and posttranslational modifications may be triggered by the studied TFs. We expect that our results could contribute to a better understanding of fungal biodiversity, especially regarding the transcription regulation involved in hydrolytic enzyme expression in T. harzianum. By describing new potential targets involved in the cellulose degradation pathway, this knowledge can be used to develop genetic manipulation strategies and expand the use of T. harzianum as an enzyme producer in biotechnological industrial applications.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

AUTHOR CONTRIBUTIONS

RR: Writing—original draft, Methodology, and Conceptualization. AA: Methodology, Software, Writing—review and editing, and Formal analysis. DA: Resources, and Writing—review and editing. JF: Writing—review and editing. MH: Resources and Writing—review and editing. AS: Supervision, review and editing, and funding acquisition.

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SUPPLEMENTARY MATERIAL

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