Distribution, Function, and Evolution of a Gene Essential for Trichotheccene Toxin Biosynthesis in *Trichoderma*

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Trichotheccenes are terpenoid toxins produced by species in 10 fungal genera, including species of *Trichoderma*. The trichotheccene biosynthetic gene (*tri*) cluster typically includes the *tri5* gene, which encodes a terpene synthase that catalyzes formation of trichodiene, the parent compound of all trichotheccenes. The two *Trichoderma* species, *Trichoderma arundinaceum* and *T. brevicompactum*, that have been examined are unique in that *tri5* is located outside the *tri* cluster in a genomic region that does not include other known *tri* genes. In the current study, analysis of 35 species representing a wide range of the phylogenetic diversity of *Trichoderma* revealed that 22 species had *tri5*, but only 13 species had both *tri5* and the *tri* cluster. *tri5* was not located in the cluster in any species. Using complementation analysis of a *T. arundinaceum tri5* deletion mutant, we demonstrated that some *tri5* homologs from species that lack a *tri* cluster are functional, but others are not. Phylogenetic analyses suggest that *Trichoderma tri5* was under positive selection following its divergence from homologs in other fungi but before *Trichoderma* species began diverging from one another. We propose two models to explain these diverse observations. One model proposes that the location of *tri5* outside the *tri* cluster resulted from loss of *tri5* from the cluster in an ancestral species followed by reacquisition via horizontal transfer. The other model proposes that in species that have a functional *tri5* but lack the *tri* cluster, trichodiene production provides a competitive advantage.

**Keywords:** *Trichoderma*, trichotheccenes, *tri5* gene, positive selection, gene deletion and complementation, antifungal activity, genomics, phylogeny

**INTRODUCTION**

Trichotheccenes are toxins produced by a wide range of fungal species from three classes of the phylum Ascomycota (Proctor et al., 2020). However, most known trichotheccene-producing fungi are members of class Sordariomycetes, order Hypocreales. These fungi include species of *Fusarium*, *Myrothecium*, *Stachybotrys*, and *Trichoderma*. Trichotheccenes are of concern because of their toxicity...
and frequent occurrence in food and feed crops pose health risks to humans and domesticated animals (Desjardins, 2006). In addition, trichothecenes can serve as virulence factors in some crop diseases caused by Fusarium species (Desjardins et al., 1996). Finally, trichothecenes have been implicated in human diseases related to fungal contamination in damp buildings (Straus, 2009).

Over 200 trichothecene analogs have been reported (Proctor et al., 2018, 2020). All the analogs share a core chemical structure, 12,13-epoxytrichothec-9-ene (EPT) and differ from one another in the types and patterns of substitutions at various positions of EPT. Trichothecenes have been classified as macrocyclic or simple, based on the presence and absence of a macrolide ring formed by a 12 or 14-atom chain esterified to EPT via hydroxyl groups at carbon atoms 4 and 15. Macrocyclic trichothecenes have this macrolide ring, whereas simple trichothecenes do not (Proctor et al., 2018).

The genus Trichoderma includes fungi that can colonize a wide variety of substrates under diverse environmental conditions. These fungi can survive as saprobes on plant debris and, in some cases, can colonize living plant tissues (Morán-Diez et al., 2009). The ability of Trichoderma species to survive and compete in such diverse habitats has been attributed in part to their collective ability to produce diverse secondary metabolites, including antimicrobial compounds that can enhance their antagonistic activity (Sivasithamparam and Ghisalberti, 1998). Since 2005, production of at least eight trichothecene analogs has been reported in multiple Trichoderma species, although not all species were reported to produce all analogs (Lee et al., 2005; Nielsen et al., 2005; Degenkolb et al., 2008; Cardoza et al., 2011; Sun et al., 2016; Ryu et al., 2017; Chen et al., 2018). This production is intriguing, because some Trichoderma species are biological control agents for crop diseases caused by other fungi (Harman, 2000; Monte, 2001; Howell, 2003; Harman et al., 2004). The biocontrol activity of some trichothecene-producing Trichoderma species may result from induction of plant defense responses in addition to antifungal activity of certain trichothecene analogs against pathogenic fungi (Malmierca et al., 2013; Gutiérrez et al., 2020).

In addition, a trichothecene analog produced by T. albolutescens was reported to have antiviral activity that contributed to protection against the Pepper Mottle Virus (Ryu et al., 2017). In trichothecene-producing fungi that have been examined, the tri5 gene encodes the enzyme trichodiene synthase, which catalyzes the cyclization of farnesyl diphosphate to trichodiene, the first committed intermediate in trichothecene biosynthesis. In most trichothecene-producing fungi, tri5 is located in a trichothecene biosynthetic (tri) gene cluster (Lindo et al., 2018; Proctor et al., 2018). However, in Trichoderma arundinaceum and T. brevicaespactum, the only two Trichoderma species whose tri clusters have been examined, tri5 is located outside the cluster in a region that does not include other known tri genes (Cardoza et al., 2011; Lindo et al., 2018; Proctor et al., 2018).

The increasing number of reports on trichothecene production in Trichoderma species suggests that production could occur widely in the genus. Therefore, we used genome sequence, biochemical, molecular genetics, and phylogenetic analyses to further characterize trichothecene production in a collection of 35 species that represent a wide range of phylogenetic diversity that exists within Trichoderma. We focused on tri5 because of its essential role in trichothecene biosynthesis combined with the uniqueness of its location outside the tri cluster in T. arundinaceum, T. brevicompactum and potentially other Trichoderma species (Cardoza et al., 2011; Proctor et al., 2020). Our results indicate that tri5 is distributed widely but not universally in Trichoderma, while the tri cluster and trichothecene production are less widely distributed. Together, results from diverse analyses suggest models that can explain the location of tri5 outside the tri cluster and why some Trichoderma species have retained a functional tri5 in the absence of the cluster.

### MATERIALS AND METHODS

#### Nomenclature

Different multispecies lineages within Trichoderma have been designated using the terms “clade,” “section,” and “core group.” Most of the lineages have been assigned names: e.g., Brevicompactum Clade, Section Trichoderma, Pachybasium Core Group (Kubíček et al., 2019). However, all these subgeneric groups are phylogenetically distinct lineages within Trichoderma. For the purposes of this study, therefore, we have used a common term, lineage, to refer to multispecies subgeneric groups within Trichoderma. In addition, multiple formats of genetic nomenclature have been used for trichothecene biosynthetic and housekeeping genes from various fungi. For consistency within the current study, we use Trichoderma genetic nomenclature for all fungal species (e.g., tri5 indicates a wild-type gene, and TRI5 indicates a protein).

#### Strains Used and Growth Conditions

For this study, we selected 35 Trichoderma species (Table 1). The species were selected based on availability of their genome sequences, previous reports of their ability to produce trichothecenes, their close relationships to known trichothecene-producing species, and/or their representation of multispecies lineages of the genus Trichoderma (Kubíček et al., 2019).

All Trichoderma strains that were examined in laboratory experiments were sporulated on CMD medium (Yellow cornmeal 1g/L, potato dextrose broth 5g/L, Agar 1.5g/L), by incubation at 28°C for 10 days, except for the strains of T. baelearicum, T. calamagrostidis, T. crystalligenum, T. psychrophilum, T. rhododendri and T. rubi, which were incubated at room temperature (15–21°C) for longer periods (3–6 weeks). Rhizoctonia solani strain ULE-R43, a fungal phytopathogen, was obtained from the University of Leon culture collection, and was used for antifungal assays on cellophane membranes. This strain was grown on potato dextrose agar medium (PDA) an incubated for 7 days at 28°C in the dark.

#### Nucleic Acid Extraction and Purification

**Growth of Fungal Strains for Genomic DNA and RNA Purification**

a. For genomic DNA extraction: six plugs from each strain, collected from the CMD cultures described above, were
### TABLE 1 | Trichoderma species and strains examined in the current study, and the occurrence of tri5, the tri cluster, and trichothecene production in the species/strains.

| Species                  | Strain no.  | Source of strain | tri5 | tri cluster | Trichothecene production | Reference               |
|--------------------------|-------------|------------------|------|-------------|--------------------------|-------------------------|
|                          |             |                  |      |             |                          |                         |
| **Lineage**              |             |                  |      |             |                          |                         |
| Brevicompactum           |             |                  |      |             |                          |                         |
| T. asperellum            | CBS 433.97  | na               | +    | −           | None                     | Vicente et al., 2020; Current study |
| T. atroviride            | IMI 206040  | −                 | −    | −           | None                     | N/A                     |
| Trichoderma              |             |                  |      |             |                          |                         |
| T. gamsii                | TO65        | ULE              | +    | −           | ND                       | Current study            |
| T. hamatum               | GD12        | na               | −    | −           | None                     | N/A                     |
| T. koningiopsis          | POS7        | na               | −    | −           | None                     | N/A                     |
| Pachybasium              |             |                  |      |             |                          |                         |
| T. polysporum            | CBS 111723  | WFBI             | +    | −           | None                     | Current study            |
| Rubi                     |             |                  |      |             |                          |                         |
| T. rubi                  | CBS 127380  | WFBI             | +    | +           | ND                       | Current study            |
| T. taxii                 | TUCIM 2377  | LID              | +    | +           | trichodermin             | Current study            |
| Deliquescens             |             |                  |      |             |                          |                         |
| T. deliquescens          | CBS 130572  | WFBI             | −    | −           | ND                       | Current study            |
| Psychrophila             |             |                  |      |             |                          |                         |
| T. baelearicum           | CBS 133222  | WFBI             | +    | +           | roridin E                | Current study            |
| T. callistosporidis      | CBS 121133  | WFBI             | +    | +           | ND                       | Current study            |
| T. crystallearum         | S38         | LWJ              | +    | +           | trichodermin, roridin A, E | Current study            |
| T. oligosporum           | CGMCC 3.17527 | na         | +    | +           | 4-acetyltrichothecolone   | Current study            |
| T. psychrophilum         | S647        | LWJ              | +    | +           | 4-acetyltrichothecolone   | Current study            |
| T. rhododendron          | CBS 119288  | WFBI             | +    | −           | ND                       | Current study            |
| Hypoccreanum             |             |                  |      |             |                          |                         |
| T. decipiens             | CBS 132861  | WFBI             | +    | −           | None                     | Current study            |
| Not assigned             |             |                  |      |             |                          |                         |
| T. alboluteosporus       | CBS 119286  | WFBI             | +    | +           | trichodermin, 16-hydroxytrichodermin | Ryu et al., 2017       |
| Semicribae               |             |                  |      |             |                          |                         |
| T. cf. fertile           | CBS 137003  | WFBI             | +    | −           | ND                       | Current study            |
| Harzianum/Virens         |             |                  |      |             |                          |                         |
| T. atrobrunneum          | ITEM 908    | na               | −    | −           | None                     | N/A                     |
| T. guizhouense          | NJAU 4742   | na              | −    | +           | None                     | Vicente et al., 2020; Current study |
| T. harzianum            | CBS 226.95  | na               | −    | −           | None                     | N/A                     |
| T. lentiforme            | CFAM-422    | na               | −    | −           | None                     | N/A                     |
| T. virens               | GV29-8      | na               | −    | −           | None                     | N/A                     |
| Sormaticum              |             |                  |      |             |                          |                         |
| T. citrinoviride         | TUCIM 6016  | na               | −    | −           | None                     | N/A                     |
| T. koningii             | JCM 1883    | na               | −    | −           | None                     | N/A                     |
| T. parareesei           | CBS125925   | na               | −    | −           | None                     | N/A                     |
| T. reesei               | QM6a        | −               | −    | −           | None                     | N/A                     |
| Longibrachiatum         |             |                  |      |             |                          |                         |
| T. cineroviride          | TUCIM 6016  | na               | −    | −           | None                     | N/A                     |
| T. kovingii             | JCM 1883    | na               | −    | −           | None                     | N/A                     |
| T. parareesei           | CBS125925   | na               | −    | −           | None                     | N/A                     |
| T. reesei               | QM6a        | −               | −    | −           | None                     | N/A                     |

*a* Subgeneric lineages of Trichoderma using names of Clades, Core Group and Sections described by Kubicek et al. (2019). The lineage Rubi was reported for the first time in this study.

*b* Sources of strains are indicated only for strains that were analyzed in laboratory experiments. DTU indicates Danish Technical University; LID indicates laboratory of Irina Druzhinina, TU Wien, Vienna, Austria; LWJ indicates the laboratory of Walter M. Jaklitsch at the University of Vienna, Vienna, Austria; ULE indicates the laboratory of Pedro A. Casquero and Santiago Gutiérrez at the University of León, Spain; WFBI indicates the Westerdijk Fungal Biodiversity Institute, Utrecht, The Netherlands; and na indicates that we examined only the genome sequence of the strain but did not acquire the strain for use in laboratory experiments.

*c* The symbol ‘+’ indicates genes that were detected in a genome sequence by BLASTn analysis, and the symbol ‘−’ indicates genes that were not detected in BLASTn analysis.

*d* ND indicates that no trichothecenes were detected using one or more of the analytical protocols described in the Methods section. None indicates that we inferred that the strain did not produce trichothecene based on the absence of tri5 and/or other known tri genes in the genome sequence. However, trichothecene production was not assessed using analytical chemical methods.

*e* Publications in which trichothecene production was determined by analytical chemical methods. N/A indicates not applicable because trichothecene production has not been assessed by chemical analysis as far as we are aware.

*Species whose genome sequences were generated during the course of the current study. All other genome sequences were downloaded from the GenBank database (Supplementary Table S1a), except the sequences for T. arundinaceum, T. brevicompactum and T. oligosporum, which were generated as part of previous studies (Chen et al., 2018; Proctor et al., 2019).*
inoculated in 100 ml YPD medium (500 ml flasks), and incubated for 24 h at 28°C and 200 rpm, except in the case of *T. balearicum*, *T. calamagrostidis*, *T. crystalligenum*, *T. psychrophilum*, *T. rhododendri*, and *T. rubi*, grown at 17°C, without shaking for 5–15 days depending on the strain.

b. For RNA extraction strains were grown following a similar procedure to that described above: six CMD plugs from each strain were inoculated in 100 ml YPD medium and incubated for 24 h at 28°C and 200 rpm. However, in the case of *T. balearicum*, due to its low growth rate and to the specific conditions required for its growth, the CMD plugs were incubated on YPD for 24 h at 28°C without shaking and later grown for 8 additional days at 17°C without shaking.

c. Wild-type *T. arundinaceum* and strains derived from it were grown as described by Lindo et al. (2018) for RNA isolation. Briefly, 5×10⁷ spores from each strain were inoculated in 50 ml of CM broth (0.5% malt extract, 0.5% yeast extract, AND 0.5% glucose) in 250-ml flasks, and incubated for 24 h at 28°C with shaking at 250 rpm. Ten milliliters of this pre-inoculum were transferred to 250 ml flasks containing 50 ml potato dextrose broth (PDB) medium and grown at 28°C with shaking at 250 rpm. After 48 h of incubation, mycelia were recovered by filtration through sterile Miracloth filters (Calbiochem, San Diego, CA), washed with 0.9% NaCl, freeze-dried, and used for DNA and/or RNA extraction.

Nucleic Acids Purification, and cDNA Synthesis
DNAs and RNAs were extracted from 20 mg of grounded mycelia using the DNeasy Plant Mini Kit (QIAGEN, Hilden, Germany) and TRIZOL reagent (InVitrogen, Carlsbad, CA), respectively, as described previously (Lindo et al., 2019). After the isolation protocol, RNA was treated with RNase-free DNase and purified further through a Zymo-Spin column (Zymo Research, Irvine, CA). cDNA synthesis was carried out using the iScript™ cDNA synthesis kit (Bio-Rad, Hercules, CA) following the manufacturer’s instructions.

Genome Sequences
Genome sequences from *T. arundinaceum*, *T. asperlum*, *T. atrobrunneum*, *T. atroviride*, *T. brevicompactum*, *T. citrinoviride*, *T. guizhouense*, *T. hamatum*, *T. harzianum*, *T. koningii*, *T. koningiopsis*, *T. lentiforme*, *T. parareesei*, *T. reesei*, *T. virens*, Aspergillus hancockii, Beauveria bassiana, Cordyceps confragosa, Fusarium fasciculatum (=phylospecies FIESC 12), Fusarium graminearum, Fusarium longipes, Microcyclospora tardicrescens, Myrothecium roridum, Spicellum ovalisporum, Stachybotrys chartarum, Stachybotrys chlorohalonata, and Trichothecium roseum were downloaded from the National Center for Biotechnology Information (NCBI) database (Supplementary Table S1). The genome sequences from the other 19 *Trichoderma* spp. examined were generated as a part of the current study using three different strategies. Strategy 1 was used for *T. albolutescens*, *T. aurantioceffusum*, *T. crystalligenum*, *T. decipiens*, *T. deliquescentes*, *T. cf. fertile*, *T. margaretense*, *T. polysporum*, *T. protrudens*, *T. psychrophilum*, *T. rodmanii*, *T. stromaticum*, *T. taxi*, and *T. turrialbense*. For these species, genomic sequences were generated at USDA with a MiSeq Illumina platform (Illumina, Inc.), and were assembled with CLC NGS Cell v. 9.5 (Qiagen, Redwood City, CA) as previously described (Proctor et al., 2018). Strategy 2 was used for *T. calamagrostidis*, *T. rhododendri*, and *T. rubi*. For these species, genome sequences were generated by the company Macrogen (Seoul, Korea; https://dna.macrogen.com) using an Illumina platform, and sequences assembly was carried out by the SPAdes (v3.15.0) assembler (Bankevich et al., 2012). Strategy 3 was used for *T. gamsii* T065 and *T. balearicum*. For these species, genome sequences were generated by the company MicrobesNG (University of Birmingham, United Kingdom; https://www.microbesng.com) using similar procedures to those included in the strategy 2. For DNA extraction, strains whose genome was sequenced using strategies 2 or 3 were grown as described above. In the case of genomes sequenced following strategy 1, strains were grown as previously described (Proctor et al., 2018).

Genomes were annotated using the gene prediction software AUGUSTUS (Hoff and Stanke, 2013). Putative secondary metabolite biosynthetic gene clusters in the genomes were identified using the software antiSMASH 6.0 (Blin et al., 2021).

Phylogenetic Analysis
Species Phylogenetic Tree
Nucleotide sequences from 20 *Trichoderma* housekeeping (=HK) genes (Supplementary Table S2) retrieved from the genome of 35 *Trichoderma* species (Table 1; Supplementary Table S1a) were used to infer a *Trichoderma* species tree. This study was carried out using two different methods, the results of which were combined. In Method 1, nucleotide sequences of the 20 HK genes from the *Trichoderma* species used in this work were retrieved as indicated above. Before the analysis, the introns from all genes were manually removed in order to have a unique continuous open reading frame in all of them. The sequences of each gene from all the *Trichoderma* spp. analyzed were individually aligned by MUSCLE software as implemented in MEGAX (Kumar et al., 2018), and then the alignments were concatenated using Sequence Matrix software (Vaidya et al., 2011). The resulting concatenated alignment was then subjected to maximum likelihood (ML) analysis as implemented in the program IQ-Tree version 1.6.7 (Nguyen et al., 2014). A second concatenated-partitioned tree was constructed by selecting for each gene the best-fit evolutionary nucleotide model deduced from the previous IQ-Tree analysis. Finally, both concatenated (non-partitioned and partitioned) alignments were subjected to ML analysis as implemented in IQ-Tree. Branch support was determined by bootstrap analysis using 1,000 pseudoreplicates. In Method 2, nucleotide sequences of each HK gene were aligned and then subjected to a ML analysis as indicated above. A consensus tree was then generated from the 20 individual HK trees as previously described (Stamatakis, 2014). Branch support in the consensus tree was determined by internode certainly (IC) as implemented in the program RAxML (Salichos et al., 2014).
**trii Evolutionary Analysis**

The presence or absence of trii in the genome sequences of 35 *Trichoderma* species was determined by BLASTn analysis (Altschul et al., 1990) using the DNA sequence of *T. arundinaceum* trii as a query. BLAST hits that were >80% identical to the query sequence were then aligned to previously described trii sequences from species of *T. arundinaceum*, also from 11 non-*Trichoderma* species in the same order as *Trichoderma* (i.e., order Hypocreales), and from two outgroup species, *Microcyclospora tardicrescens* (order Capnodiales, class Dothideomycetes) and *Aspergillus hancockii* (order Eurotiales, class Eurotiomycetes), and a phylogenetic tree was then inferred from the alignment (Proctor et al., 2020). trii sequences were aligned by the CLUSTAL software as implemented in MEGA7 (Kumar et al., 2016) and intron sequences were manually removed. The resulting ORFs were translated to amino acid sequences, aligned, and then converted back to the original nucleotide sequences before further analysis. Aligned trii sequences were subjected to maximum likelihood (ML) analysis, and the best substitution model was determined with MEGA7 under the Akaike Information Criterion (AIC). The best model was GTR (General Time Reversible) with a discrete gamma distribution and an allowance for the presence of invariant sites (Nei and Kumar, 2000). Bootstrap analysis with 1,000 pseudoreplicates was used to determine the significance of nodes.

For assessments of horizontal gene transfer and selection, a tree was also inferred from the concatenated alignment of a subset of six housekeeping genes (*cpr1, dpa1, fas1, fas2, lcb2*, and *rpb1; Supplementary Table S2*) using the same procedure described above to infer the species tree. However, to avoid unwanted sources of variation, we used the same nucleotide substitution model that was used to infer the trii tree to infer the six-housekeeping gene (6HK) tree. Branch support in the 6HK tree was assessed by both bootstrap and internode certainly analyses as described above.

**Estimation of Divergence of trii**

To assess divergence and positive selection of trii we compared estimates of the number of changes in codon sequences that change amino acid sequence (i.e., the number of nonsynonymous substitutions per nonsynonymous site, or dN) and changes in codon sequences that do not change amino acid sequence (i.e., synonymous substitutions per synonymous site, or dS). The estimates were obtained using the modified Nei-Gojobori model (Zhang et al., 1998) as implemented in MEGA7 (Kumar et al., 2016). We then estimated selection of trii by determining the dN/dS ratio (ω) for pairwise combinations of species of (i) *Trichoderma* versus *Trichoderma* (TT), (ii) non-*Trichoderma* versus non-*Trichoderma* (NN), and (iii) *Trichoderma* versus non-*Trichoderma* (TN). ω values and statistics were obtained using the R statistics software package.

Positive selection was also assessed using CodeML as implemented in the program package PAML4 (Yang, 2007). Two models were compared with CodeML: (i) a null hypothesis (H0) in which ω was homogeneous in all branches in the trii tree, and (ii) an alternative hypothesis (H1), in which ω differed in each branch in the *Trichoderma* clade of the trii tree. The possible differences between these two models were statistically analyzed with the Kishino–Hasegawa test (Kishino and Hasegawa, 1989). The program BaseML, also included in PAML, was used to compare the different tree topologies versus a multiple alignment. Finally, the Shimodaira–Hasegawa test was used to determine if the topologies of the trii and housekeeping gene trees were statistically different (pSH < 0.05; Shimodaira and Hasegawa, 1999).

**Trichothecene Detection and Quantification**

Trichothecene analogs and pathway intermediates were analyzed using three analytical systems: HPLC-UV, GC-MS, and LC-MS. The HPLC-UV system was used to detect and quantify harzianum A (HA) and consisted of a high performance liquid chromatography fitted with Waters YMC analytical column (150 by 4.6 mm) and coupled to ultraviolet light absorption detector as previously described (Cardoza et al., 2011). The GC-MS system was used to detect and, in some cases, quantify all other simple trichothecenes and consisted of Hewlett Packard 6890 gas chromatograph fitted with a HP-5MS column (30 m length, 0.25 mm film thickness) coupled to an Agilent 5,793 mass detector as described previously (Lindo et al., 2018). The LC-MS system was used to detect macrocyclic trichothecenes and consisted of a Thermo Dionex Ultimate 3000 liquid chromatograph fitted with a Phenomenex Kinetic F5 column (150 mm length, 2.1 mm diameter, 1.7 μm particle size) coupled to the electrospray interface of a Thermo QExactive mass spectrometer operated in positive mode. The procedures for chromatographic separation and identification of trichothecene analogs were adapted from those previously described (Proctor et al., 2018). Together, the GC-MS, LC-MS, and HPLC systems used in this study had the capacity to detect over 150 trichothecene analogs (Cole et al., 1981; Savard and Blackwell, 1994; Lee et al., 2005; Nielsen et al., 2005; Degenkolb et al., 2008; Cardoza et al., 2011; Sun et al., 2016; Ryu et al., 2017; Chen et al., 2018).

For HPLC-UV analysis, strains were grown using the conditions described for RNA isolation described above. Filtrates from the resulting cultures were extracted with ethyl acetate. For the GC-MS and LC-MS analyses, strains were grown in liquid YEPD medium (0.1% yeast extract, 0.1% peptone, and 2% glucose) for 7 days after which the resulting cultures (growth and culture medium) were extracted with ethyl acetate. All other experimental parameters (e.g., ratio of solvent volume to culture volume, injection volume, and chromatographic mobile phases) have been previously described (Cardoza et al., 2011; Proctor et al., 2018).

**Plasmid Construction**

**Construction of pαtri5**

1,180-bp and 1,061-bp fragments, corresponding to the 5'- and 3'-flanking regions to the T. arundinaceum trii gene were amplified by PCR using the Q5 high-fidelity DNA polymerase (New England Biolabs, County Road, MA) and oligonucleotides trii_5rF_BamHI/
tri5_5r_Smal and tri5_3rF_Smal/tri5_3r_SalI, respectively (Supplementary Table S3). The amplicon corresponding to the 3’-flanking region was subcloned in pBluescript II KS+ (Stratagene, La Jolla, CA) previously digested with EcoRV, and dephosphorylated with alkaline phosphatase (Fermentas, Vilnius, Lithuania). The resulting plasmid, pBT5_3R (4,022 bp) was digested with BamH1/Smal and ligated to the tri5-5’-flanking amplicon, previously digested with the same endonuclease enzymes, to originate the plasmid pBT5_3R-5R (5,169 bp). The resulting plasmid was linearized with Smal, dephosphorylated, and then ligated to the hygR (hygromycin resistance) cassette (2,708 bp), consisting of the coding region of the hygromycin phosphotransferase gene (hph) from E. coli, fused to the gpdA promoter and trpC terminator sequences of A. nidulans. HygR cassette was released from plasmid pAN71 (Punt et al., 1987) by HindIII digestion, treatment with Klenow fragment of DNA polymerase I (Fermentas), and a final digestion with Ecl136II. The resulting plasmid pATri5 (7,877 bp) was used to delete tri5 coding region of T. arundinaceum by a double cross-over strategy (Casqueiro et al., 1999).

Construction of Plasmids to Express the tri5 Gene From Different Trichoderma Species Into the TARUN tri5 Gene Deleted Mutant (Δtri5.3)

The tri5 ORFs were amplified by PCR using the Q5 high-fidelity DNA polymerase (New England Biolabs) from genomic DNAs of T. arundinaceum (1,226 bp), T. baealicum (1,243 bp), T. decipiens (1,176 bp), T. cf. fertile (1,142 bp), T. gamsii TO65 (1,227 bp), T. polysporum (1,223 bp), and T. stromaticum (1,157 bp), and 5’-phosphorylated oligonucleotides TARUN_T5_5/TARUN_T5_3, TBALE_T5_5/TBALE_T5_3, TDECI_T5_5/TDECI_T5_3, TERT_T5_5/TERT_T5_3, TGAMS_T5_5/TGAMS_T5_3, TPOLY_T5_5/TPOLY_T5_3, and TSTRO_T5_5/TSTRO_T5_3, respectively (Supplementary Table S3). The amplicons were subcloned in plasmid pTACbh (Cardoza et al., 2015) previously digested with NcoI, filled with Klenow, and dephosphorylated. The resulting plasmids were linearized with EcoRI, except in the case of pTACbh containing T. arundinaceum tri5 that was digested with HindIII. Then, plasmids were filled with klenow, dephosphorylated, and ligated to the 1,591-bp bleR (bleomycin/pleomycin resistance) cassette that was isolated and purified from plasmid pJL43b1 (Gutiérrez et al., 1997), by digestion with HindIII, filled with klenow, and again digested with Ecl136II. The final plasmids [pTCT_TSTTARUN_ble (8,164 bp), pTCT_TSTTBALE_ble (8,181 bp), pTCT_TSTDECI_ble (8,114 bp), pTCT_TSTTERT_ble (8,080 bp), pTCT_TSTGAMS_ble (8,165 bp), pTCT_TSTPOLY_ble (8,161 bp), pTCT_TSTSTRO_ble (8,095 bp)] (Supplementary Figure S1b-h) were linearized with the endonuclease NdeI to transform protoplasts of the Δtri5-3 mutant. The tri5 genes from the different Trichoderma strains used in this study, once subcloned as described above, were fully sequenced to confirm their integrity.

Trichoderma Transformation and Selection of Transformants

Transformation of T. arundinaceum to obtain tri5-deletion mutants and transformants for heterologous expression of tri5 homologs was carried out using a protoplast-mediated protocol as previously described (Proctor et al., 1999; Cardoza et al., 2006). Selection of tri5-deleted and tri5-complemented transformants was carried out on a regeneration medium containing hygromycin 150 μg/ml or phleomycin 100 μg/ml, respectively (Malmierca et al., 2012; Cardoza et al., 2019).

Generation of a T. arundinaceum IBT 40837 tri5-Deleted Mutant (Δtri5)

Deletion of tri5 was achieved by transforming protoplasts of the T. arundinaceum strain IBT 40837 with plasmid pATri5 (7,877 bp, Supplementary Figure S1a) that had been linearized with XhoI endonuclease. Colonies growing on the hygromycin B-amended regeneration medium were analyzed by the Terra PCR method (Takara Bio.) using two oligonucleotide primer pairs designed to detect homologous recombination between the deletion construct and the 5’ (primer pair T5-5rr and TtrpC-d) and 3’ (primer pair Pgpda-d and T5-3rr) flanking regions of tri5 (Supplementary Table S3). Transformants that yielded both amplicons were also analyzed by PCR for the presence of a 674-bp fragment internal to T. arundinaceum tri5. Transformants that yielded the 674-bp amplicon were then subject to Sanger sequence analysis of the T5-5rr/TtrpC-d and Pgpda-d/T5-3rr amplicons to confirm that the amplicons originated from the expected double cross-over event. One transformant from which the expected sequence data for the 5’ and 3’ flanking regions was obtained was selected for genome sequence analysis, following procedure #1 described above, to further confirm deletion of the tri5 coding region.

Real Time qPCR

For qPCR analysis primers for tri5 and actin-encoding genes were designed for all Trichoderma species analyzed in the present work (Supplementary Table S3). The primer pairs amplification efficiencies range between 91.3 and 110.6% (Supplementary Table S3). cDNAs were quantified using a Nanodrop ND-1000 (ThermoFisher), and qPCR reactions were carried out on a Step One system (Applied Biosystems) using two oligonucleotide primer pairs designed to detect homologous recombination between the deletion construct and the 5’ (primer pair T5-5rr and TtrpC-d) and 3’ (primer pair Pgpda-d and T5-3rr) flanking regions of tri5 (Supplementary Table S3). Transformants that yielded both amplicons were also analyzed by PCR for the presence of a 674-bp fragment internal to T. arundinaceum tri5. Transformants that yielded the 674-bp amplicon were then subject to Sanger sequence analysis of the T5-5rr/TtrpC-d and Pgpda-d/T5-3rr amplicons to confirm that the amplicons originated from the expected double cross-over event. One transformant from which the expected sequence data for the 5’ and 3’ flanking regions was obtained was selected for genome sequence analysis, following procedure #1 described above, to further confirm deletion of the tri5 coding region.

Volatile Collection and Characterization of Terpene Compounds Emitted From Trichoderma Cultures

Volatile compounds emitted from the transformants expressing the tri5 from T. cf. fertile and T. gamsii TO65 were collected using the previously described closed-loop stripping method (Laraba et al., 2020). Briefly, a 7-days old culture grown on potato dextrose agar (PDA) Petri dish (60 mm × 15 mm) was sealed in a 3L glass desiccator. The air space within the desiccator was recirculated through a filter containing 25 mg of Porapak™ Porous Polymer adsorbent, type Q, 80–100 mesh (Supelco, Bellefonte, PA, United States) for 24 h at a rate of December 2021 | Volume 12 | Article 791641

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3 volumes of total air space per hour. The compounds were then eluted from the adsorbent with 150 μl of dichloromethane and 2 μl of the effluent were analyzed on an Agilent 6890 chromatograph/Agilent 5973 mass spectrometer. The compounds were separated on a HP-5MS column held at 50°C for 3 min after injection, and then the temperature was gradually heated to 250°C at 30°C/min where it was held for 1 min. Individual peaks were identified based on comparison of ion fragmentation patterns. The terpene compounds were initially identified based on NIST 11 library spectral matches and then verified with standards.

**Antifungal Assays on Cellophane Membranes**

These assays were performed as described previously (Cardoza et al., 2015), but incubating the Trichoderma plugs over the cellophane membranes for only 24 h instead of the 48 h previously recommended. Rhizoctonia solani ULE-R43 was the pathogen used in these assays, and its growth was followed until the pathogen’s mycelium covers all the surface in the control plates, i.e., plates with the pathogen growing without previous growth of Trichoderma, which usually took 6–7 days at 28°C in the dark. Three biological replicates were used for each Trichoderma strain.

**RESULTS**

**Phylogeny of the Trichoderma Species**

Together, the 35 Trichoderma species selected for this study represented 10 previously described multispecies lineages and three species (T. alboluteescens, T. rubi and T. taxi) that were not previously assigned to lineages (Table 1). The previously described lineages Brevicom pactum, Longibrachiatum, Harzianum/Virens, Psychrophila, Trichoderma were represented by 2–7 species each, while the previously described lineages Deliquescens, Hypocreanum, Pachybasium, Seniorbis, Stromaticum were represented by only one species each. To better understand the phylogenetic relationships of all 35 species, we inferred a species tree by maximum likelihood analysis of concatenated alignments of 20 housekeeping genes that were retrieved from genome sequences of the fungi (Figure 1). In the resulting tree, the five previously described lineages represented by multiple species were resolved into well supported and exclusive clades with bootstrap values of 100 and internode certainty values of 0.69–1.0 (Figure 1). For most other branches in the species tree, bootstrap values were 100 and internode certainty values ranged from 0.19 to 0.69. Only one branch had a bootstrap value less than 70: the branch that included the Brevicom pactum, Trichoderma, and Pachybasium lineages as well as T. rubi and T. taxi (Figure 1). T. rubi and T. taxi were resolved as a well-supported clade, which we hereafter refer to as the Rubi lineage. The topology of the phylogenetic tree (Figure 1) was largely consistent with multiple species phylogenies that have been previously reported for other combinations of Trichoderma species (Jaklitsch and Voglmayr, 2015; Kubicek et al., 2019). Given this and the high levels of branch support, we used the species tree in Figure 1 to provide a phylogenetic context for Trichoderma species and lineages in subsequent analyses.

**Distribution of tri5 Gene in Selected Trichoderma Species**

We used BLASTn analysis of genome sequences as an initial screen for the presence of tri5 in the 35 Trichoderma species. This analysis indicated that a single copy of tri5 was present in 22 species and absent in the other 13 species (Table 1; Figure 2). Of the 22 species with tri5, multiple similarities in the content and arrangement of flanking genes indicate that tri5 is in the same genomic location, which we designated Genomic Region 1 (GR1), in 18 species and different genomic locations in each of four other species: GR2 in T. asperellum; GR3 in T. gamsii T065; GR4 in T. polysporum; and GR5 in T. stromaticum. Although the latter four species had tri5, results of BLASTn analysis indicated that they do not have any other known tri genes. Further, in all 22 species with tri5, the tri5 flanking genes did not share significant sequence homology with known tri genes according to BLASTn and BLASTx analyses. This latter finding is consistent with the physical separation of tri5 and the tri cluster that was previously reported for T. arundinaceum and T. brevicompactum (Cardoza et al., 2011; Proctor et al., 2018). The degree of conservation of genes in the tri5-flanking region was higher among species from the same lineage than those from different lineages (Figure 2). Based on sequence similarities to proteins with known functions, proteins encoded by some tri5-flanking genes included a cytochrome P450 monooxygenase, AraC-type transcriptional factor, major facilitator superfamily (MFS) transporter, homoserine acetyl transferase, oligosaccharidyl lipid flipase, and ATP-dependent RNA helicase, suggesting that some of the flanking genes could be involved in secondary metabolite biosynthesis while others are unlikely to be involved (Figure 2; Supplementary Table S5).

**Trichothecene Production**

Our assessment of trichothecene production by Trichoderma species was limited by the availability of strains. Nevertheless, based on analyses in the current study and previously published reports, 11 out of the 35 species produced trichothecenes (Figure 3; Supplementary Figure S2; Table 1). Producing species were from the multispecies lineages Brevicom pactum and Psychrophila as well as T. alboluteescens and T. taxi. Of the seven species examined in the Brevicom pactum lineage, five produced trichothecenes (Table 1). These five species produced one or more of the following simple trichothecene analogs: trichodermol, trichodermin, isotrichodermin, and harzianum A. Of the six species examined from the Psychrophila lineage, four produced trichothecenes. These species produced simple (e.g., trichodermin) or macrocyclic (e.g., roridins A, D, and/or E) trichothecenes or a mixture of both types of trichothecenes (Figure 3; Table 1). T. taxi produced trichodermin. Our analysis did not detect trichothecene production in T. alboluteescens strain CBS 119286, which originated in Germany,
but previous report indicated that another strain from Korea can produce trichodermin and 16-hydroxytrichodermin (Table 1; Ryu et al., 2017). We had genome sequences but not cultures for 11 Trichoderma species/strains and, therefore, could not analyze their trichothecene production profile. However, based on the absence of tri5 and/or other tri
FIGURE 2 | Analysis of the gene content in homologs of the *T. arundinaceum* trnS region (GR1) in 35 *Trichoderma* species. trnS was absent in GR1 homologs from 17 species and occurred at genomic locations other than GR1 in four species: GR2 in *T. asperellum*, GR3 in *T. gamsii* T065, GR4 in *T. polysporum*, and GR5 in *T. stromaticum*. Genes are represented by arrows that point in the direction of transcription. Numbers or letters above genes correspond to the numbers in the list of 37 proteins on the right side of the figure; the list indicates the predicted gene/protein functions based on sequence homology. Genes that are potentially involved in secondary metabolite biosynthesis and/or included in subsequent analyses are depicted as colored arrows. *trnS* is depicted as a bright green arrow; genes that do not exhibit homology to genes of known function are depicted as black arrows, and all other genes are depicted as gray arrows. Groups of arrows that are within a yellow or blue rectangle indicate multigene segments of DNA inserted into GR1. Names in red type on the left side of the figure are lineages of *Trichoderma* (Kubicek et al., 2019).

(Continued)
genes in the genome sequences, we inferred that these strains could not produce trichothecenes (Table 1).

Production of trichothecenes by all species in the Brevicompactum lineage, except T. margaretense and T. aurantioeffusum, is consistent with the presence of tri5 and other tri genes in the producing species and the absence of most or all of the genes in T. margaretense and T. aurantioeffusum (Table 1; Figure 3). In fact, when trichothecene production was initially described in Trichoderma species, it was proposed that production was limited to members of the Brevicompactum lineage (Cardoza et al., 2011). However, the results of the current study indicate that production, and presumably all necessary tri genes, occur in other lineages of the genus and occurs widely among members of in the Psychrophila lineage (Figure 2).

Variation in tri5 Genomic Region GR1
Analysis of genomic region GR1 revealed variation in the presence and absence of genes in homologs of the region in species from different Trichoderma lineages. There were two differences that stood out because they involved insertions of multigene segments of DNA into GR1.

The first multigene-segment insertion included 5–6 genes and occurred in members of the Harzianum/Virens and Stromaticum lineages (shaded in blue in Figure 2). In the segment, gene 24 was predicted to encode urea carboxylase (UC; Figure 2), which is thought to be involved in urea utilization (Navarathna et al., 2010). In the same segment, gene 26 encoded a NADPH dehydrogenase, gene 23 encoded a lactam utilization protein (LamB), gene MFS encoded a major facilitator superfamily, and gene D encoded a Zn Cys\(_2\)His\(_2\) transcription factor. Whether these genes constitute a urea utilization gene cluster is not known. Homologs of these UC-related genes were also present and located adjacent to one another in T. koningiopsis, T. gamsii T065 and T. atroviride (Trichoderma lineage) and T. polysporum (Pachybasium lineage). However, in these latter four species, the UC-related genes were not located in GR1 (Figure 2).

The second multigene-segment insertion in GR1 occurred in four of six species in the Psychrophila lineage: T. balearicum, T. calamagrostidis, T. crystalligenum, and T. oligosporum (shaded in yellow in Figure 2). In Figure 2, only four genes in this segment are shown, but an in-depth analysis revealed that the segment consisted of 15 genes (Supplementary Figure S3).
The segment also occurred in *T. psychrophylum* and *T. rhododenrī*, which are also members of the *Psychrophila* lineage, but in these latter two species the segment was not located in GR1 (Supplementary Figure S3). Whether the 15 genes constitute a cluster is not known. However, a search for homologs of the segment using the antiSmash 6.0 software (Blin et al., 2021) revealed significant synteny with the agnestin biosynthetic gene cluster in *Paecilomyces divaricatus* (Szwalbe et al., 2019; Supplementary Figure S3).

**Phylogenetic Analysis of the *tri5* Gene**

To assess the relationships of *tri5* homologs, we inferred a maximum likelihood tree from an alignment of full-length *tri5* exon sequences from 22 species of *Trichoderma* and 13 species from nine other fungal genera (Figure 4, left panel; Supplementary Table S1b). In the resulting tree, *Trichoderma tri5* sequences formed a well-supported and exclusive clade (bootstrap value = 100; Figure 4). Within the *Trichoderma* clade, bootstrap support for nodes varied from <70 to 100. The topology within the *Trichoderma* clade was consistent with the species tree (Figure 4, right panel) in some respects but not in others. For example, the *Brevicompactum* and *Psychrophila* lineages formed monophyletic clades in both trees, although bootstrap values for these clades were 78 and 89, respectively, in the *tri5* tree but 100 in the species tree (Figures 1, 4). In contrast, although members of the *Trichoderma* lineage formed a well-supported and exclusive clade in the species tree, they were not monophyletic in the *tri5* tree. The relationships of *tri5* homologs from non-*Trichoderma* species were similar to those described in previous studies (Proctor et al., 2018, 2020). Another notable feature of the *tri5* tree was the long branch separating the *Trichoderma* clade from other non-*Trichoderma* clades (indicated with a red dot in Figure 4-left panel). This branch was longer than all other branches in the *tri5* tree and longer than the equivalent branch in the species tree (Figure 4, right panel; Supplementary Figure S4).

**Analysis of *tri5* Horizontal Transfer**

Horizontal transfer of *tri5* to *Trichoderma* is one possible explanation for the branch conflicts in the species tree and the *tri5* tree as well as for the long branch leading to the *Trichoderma* clade in the *tri5* tree. In BLASTn and BLASTp analyses using *Trichoderma tri5* sequences as queries against the NCBI database, the highest-scoring hits were always *tri5* homologs from other *Trichoderma* species. In contrast, *tri5* homologs from other fungi had substantially lower BLAST scores and percent identity values. Thus, the results of BLAST analyses did not provide evidence for a potential donor in the putative horizontal transfer of *tri5* to *Trichoderma*.

To further investigate the possibility of horizontal transfer of *tri5* to *Trichoderma*, we used the Shimodaira-Hasegawa test to assess whether the *tri5* tree topology was significantly different than topologies of trees inferred from housekeeping genes. Our rationale was that if *tri5* had been horizontally transferred to *Trichoderma* from one of the other fungi included in this analysis, the Shimodaira-Hasegawa tests should indicate that the *tri5* tree topology was consistently different than the topology of all housekeeping gene trees. Therefore, in the analysis we inferred maximum likelihood trees from an alignment of *tri5*, individual alignments of six housekeeping genes, and a concatenated alignment of these six housekeeping genes. We then used the Shimodaira-Hasegawa test to determine whether the topology of each of the resulting (original) trees was more likely than topologies of trees inferred from the other alignments. The results of the analysis indicated that for each housekeeping gene alignment the original tree was more likely than the *tri5* tree topology (Table 2). The topology of the *tri5* alignment is rejected by all the HK alignments (pSH < 0.05). However, the *tri5* alignment rejects (pSH < 0.05) only the *fas1* and *lcb2* topologies, but it is compatible (pSH > 0.05) with topologies of 6HK, *cpr1*, *dpa1*, *fas2*, and *rpb1* trees (Table 2). Thus, the results of this analysis indicate that the topology of the *tri5* tree was not significantly different from the topologies of all the housekeeping gene trees, a finding that does not support the proposed horizontal transfer of *tri5*.

**Assessment of *tri5* Selection**

The greater length of the *Trichoderma* branch relative to other branches in the *tri5* tree (Figure 4) could also result from rapid divergence of ancestral *Trichoderma tri5* after it diverged from other lineages of the gene. To assess whether this apparent rapid divergence resulted from positive selection of *tri5*, we compared estimates of the number of nonsynonymous substitutions per nonsynonymous site (dN), number of synonymous substitutions per synonymous site (dS), and the dN/dS ratio (=ω) for *tri5* and six housekeeping genes in pairwise combinations of species. We consolidated the comparisons into three types: i) non-*Trichoderma* versus non-*Trichoderma* (NN), ii) *Trichoderma* versus non-*Trichoderma* (TN) and iii) *Trichoderma* versus *Trichoderma* (TT; Figure 5).

The result of this analysis revealed that dN values for *tri5* were higher in *Trichoderma* vs. non-*Trichoderma* comparisons than in non-*Trichoderma* vs. non-*Trichoderma* comparisons (Figure 5; Table 3). A similar difference was observed for ω values for *tri5* in *Trichoderma* vs. non-*Trichoderma* comparisons relative to non-*Trichoderma* vs. non-*Trichoderma* comparisons (Figure 5; Table 3). The dN and ω values in *Trichoderma* vs. non-*Trichoderma* comparisons were also higher for *tri5* than for the housekeeping genes (Figure 5; Supplementary Figure S5), suggesting a higher rate of nonsynonymous changes in *tri5* than in the housekeeping genes. In contrast, the dN and ω values for *tri5* in *Trichoderma* vs. *Trichoderma* comparisons were not significantly different than for housekeeping genes, suggesting the rate of nonsynonymous changes in *tri5* slowed in *Trichoderma* species since they began diverging from one another. In contrast to the differences in dN and ω values noted above, *tri5* and housekeeping genes dS values were similar in *Trichoderma* vs. non-*Trichoderma* and non-*Trichoderma* vs. non-*Trichoderma* comparisons (Figure 5; Table 3; Supplementary Figure S5).

To further assess divergence of *tri5*, we used the program CodeML to compare two hypotheses: a null hypothesis (Hₐ)
TABLE 2 | Probability values from a Shimodaira-Hasegawa (pSH) test that assessed whether the original maximum likelihood trees inferred from each sequence alignment was a better fit than trees inferred from other alignments.

| Topology | 6HK | cpr1 | dpa1 | fas1 | fas2 | lcb2 | rpb1 | tri5 |
|----------|-----|------|------|------|------|------|------|------|
| 6HK      | -   | 0.937| 0.866| 0.798| 0.562| 0.798| best | 0.625|
| cpr1     | 0.398| -    | best | 0.864| 0.548| 0.752| 0.300| 0.416|
| dpa1     | 0.006**| 0.214| -    | 0.100| 0.036| 0.506| 0.251| 0.107|
| fas1     | 0.000***| 0.012*| 0.001**| -    | 0.000***| 0.000***| 0.000***| 0.000***|
| fas2     | 0.875| 0.934| 0.722| best | -    | 0.676| 0.808| 0.289|
| lcb2     | 0.000***| 0.711| 0.035*| 0.027*| 0.002| -    | 0.011*| 0.034*|
| rpb1     | 0.001**| 0.518| 0.017*| 0.281| 0.001**| 0.476| -    | 0.486|
| tri5     | 0.000***| 0.000***| 0.000***| 0.000***| 0.000***| 0.000***| 0.000***| -    |

A value of pSH<0.05 indicates that a tree inferred from another alignment was a significantly worse fit than the original tree.

One asterisk (*) indicates p values of 0.01–0.05; two asterisks (**) indicates p values of 0.001–0.0099; and three asterisks indicate p values <0.001. Note that for all housekeeping gene alignments, the tri5 tree topology can be discarded (pSH<0.05) with the data of any of the HK genes, i.e., it was a worse fit than the original housekeeping gene trees. However, the best topologies, 6HK, cpr1 and fas2 tree topologies, can not be discarded (pSH >0.05) by any of the alignments.

FIGURE 4 | (Left panel) Phylogenetic tree inferred by maximum likelihood analysis of full-length exon sequences of tri5 from Trichoderma species and other fungi. Numbers on each branch are bootstrap values based on 1,000 pseudoreplicates. Blue rectangles denote species that have tri5 but no other tri genes. Red and green rectangles denote species used in the Δtri5 mutant complementation analysis; a green rectangle indicates successful complementation, and a red rectangle indicates no complementation. Yellow highlighting denotes species that were both used in the complementation analysis and in which tri5 is located at GR1. (Right panel) Phylogenetic tree inferred by maximum likelihood analysis of full-length exon sequences of six housekeeping genes (6HK; Supplementary Table S2) from the same species included in the tri5 tree. Numbers on branches are bootstrap values (blue type) based on 1,000 pseudoreplicates and internode-certainly values (red type). In both panels, lineage names, as described previously (Kubicek et al., 2019), are indicated in red type to the right of each tree. The red circle in both trees denotes the different lengths of the Trichoderma branch in the two trees.
in which $\omega$ was the same for all branches in the tri5 tree, and an alternative hypothesis ($H_0$) in which $\omega$ for the Trichoderma branch ($\omega_T$) differed from other branches ($\omega_o$) in the tri5 tree. The results of this analysis revealed that the Trichoderma branch had a significantly higher $\omega$ than other branches in the tri5 tree (Table 4). This in turn suggests positive selection within the Trichoderma branch of the tri5 tree. A caveat to this result was that in the housekeeping gene tree the Trichoderma branch had a significantly higher $\omega$ than other branches. Nevertheless, the $\omega_T/\omega_o$ ratio in the tri5 tree was substantially higher than in the housekeeping gene trees (Table 4).

Phylogenetic Analysis of Trichoderma tri5-Flanking Genes
If the long Trichoderma branch in the tri5 tree resulted from positive selection, it is possible that this selection affected neighboring genes through the phenomenon of genetic hitch-hiking (Barton, 2000). To determine whether the putative selection of tri5 affected divergence of neighboring genes, we inferred trees from homologs of three Trichoderma tri5-flanking genes that encoded a putative homoserine acetyltransferase, a oligosaccharidyl lipid flippase, and a ATP-dependent RNA helicase (Figure 2). These three genes were selected because of their proximity to tri5 in Trichoderma and the occurrence of closely related homologs in other fungi included in this study. In maximum likelihood trees inferred from the three genes, the Trichoderma branch was more similar in length to the Trichoderma branches in housekeeping gene trees than in the tri5 tree. These results suggest that the putative positive selection on the Trichoderma tri5 did not significantly affect divergence of the acetyltransferase, flippase, and helicase genes (Supplementary Figure S6).

Analysis of Functionality of Selected Trichoderma tri5 Homologs
The finding that some Trichoderma species have tri5 but no other known tri genes (Table 1) raises the question about the functionality of the tri5 in these species. Further, it is not known whether the different genomic locations of tri5 can affect its function. To address these questions, we analyzed expression of the tri5 homologs in their native locations and in a heterologous system.

Expression of tri5 in Selected Trichoderma Species
We compared expression of tri5 in the following selected Trichoderma species: i) T. arundinaceum and T. balearicum, which produce trichothecenes, have a tri cluster, and have tri5 in genomic region GR1; ii) T. cf. fertile and T. decipiens, which do not produce trichothecenes, lack the tri cluster, and have tri5 in genomic region GR1; and iii) T. gamsii T065, T. polysporum, and T. stromaticum, which do not produce trichothecenes, lack the tri cluster, and have tri5 in genomic region GR3, GR4 and GR5, respectively. The results of the analysis indicated high levels of tri5 expression in T. arundinaceum, no detectable expression in T. cf. fertile, and very low levels of expression in the other five species examined (Figure 6). Surprisingly, tri5 expression in T. balearicum was very low even though it produced the trichothecene roridin E under the same conditions (Table 1; Figure 3).
### TABLE 3 | Divergence values (mean ± sd) for pairwise combinations of sequences.

|  | crp1  | dpa1  | fas1  | fas2  | lcb2  | rpb1  | 6HK    | tri5  |
|---|-------|-------|-------|-------|-------|-------|--------|-------|
| dN |       |       |       |       |       |       |        |       |
| NN | 0.188 ± 0.083 | 0.208 ± 0.124 | 0.109 ± 0.058 | 0.136 ± 0.069 | 0.185 ± 0.104 | 0.196 ± 0.125 | 0.160 ± 0.088 | 0.187 ± 0.056 |
| TN | 0.199 ± 0.042 | 0.178 ± 0.065 | 0.111 ± 0.035 | 0.121 ± 0.045 | 0.191 ± 0.062 | 0.146 ± 0.097 | 0.143 ± 0.058 | 0.396 ± 0.030 |
| TT | 0.052 ± 0.018 | 0.029 ± 0.009 | 0.028 ± 0.009 | 0.022 ± 0.007 | 0.067 ± 0.031 | 0.011 ± 0.004 | 0.028 ± 0.009 | 0.076 ± 0.049 |
| dS |       |       |       |       |       |       |        |       |
| NN | 0.946 ± 0.274 | 1.216 ± 0.290 | 0.906 ± 0.300 | 0.927 ± 0.309 | 1.064 ± 0.325 | 1.015 ± 0.262 | 0.988 ± 0.279 | 1.048 ± 0.306 |
| TN | 0.917 ± 0.153 | 1.269 ± 0.134 | 0.963 ± 0.180 | 0.985 ± 0.17 | 1.076 ± 0.153 | 1.138 ± 0.1 | 1.053 ± 0.132 | 1.055 ± 0.198 |
| TT | 0.411 ± 0.120 | 0.567 ± 0.157 | 0.446 ± 0.128 | 0.468 ± 0.127 | 0.482 ± 0.145 | 0.48 ± 0.13 | 0.476 ± 0.131 | 0.670 ± 0.234 |

Comparisons were performed for pairs of sequences from two non-Trichoderma (NN), two Trichoderma (TT), or a Trichoderma and a non-Trichoderma (TN) species. Values are estimates of synonymous changes per synonymous site (dS); nonsynonymous changes per nonsynonymous site (dN), and the ratio of the two values (i.e., dN/dS = ω).

Alignments of six housekeeping genes were included in this analysis. The trees inferred from tri5 and the concatenated alignments of six housekeeping genes are shown in Figure 4.

*Values in bold type indicate tri5 comparisons of Trichoderma versus non-Trichoderma pairs (TN) that yielded relatively high dN and dS values.

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### Development of Heterologous tri5 Expression System

We rationalized that a tri5 deletion mutant (Δtri5) of *T. arundinaceum* would serve as an effective heterologous expression system for tri5 homologs from other *Trichoderma* species, because the mutant should not produce trichothecenes, but functional homologs of tri5 should complement the mutant to restore trichothecene production. Using the deletion protocols described in the Methods we identified three Δtri5 mutants of *T. arundinaceum* (Figure 7). None of them produced detectable levels of harzianum A under conditions that induced production of high levels of the trichothecene in the wild-type progenitor strain (Supplementary Figure S7). Genome sequence analysis of one of the mutants, strain Δtri5.3, confirmed tri5 deletion. Therefore, we selected strain Δtri5.3 as the host for heterologous expression analyses.

### Heterologous Expression of tri5 Homologs From Selected *Trichoderma* Species

For heterologous expression analysis, we selected tri5 homologs from *T. balearicum*, *T. cf. fertile*, *T. gamsii* T065, *T. decipiens*, *T. polysporum*, *T. stromaticum*, and *T. arundinaceum* (positive control). A separate tri5 overexpression plasmid was constructed for each of these tri5 homologs, and in each plasmid the tri5 coding region was fused to the *T. harzianum* Tadtr gene promoter, which confers high levels of expression in *T. arundinaceum* (Proctor et al., 2018; Cardoza et al., 2019; Lindo et al., 2019; Supplementary Figures S1b–h). Each plasmid was transformed separately into Δtri5 mutant strain Δtri5.3, and 20 of the resulting transformants recovered for each construct were analyzed by PCR to verify integration of the tri5-expression cassette. Two transformants for each construct that yielded the expected amplicons were then selected for trichothecene analysis. Transformants carrying tri5 homologs from *T. arundinaceum*, *T. balearicum*, *T. decipiens*, *T. stromaticum*, and *T. polysporum* produced harzianum A (HA), but the levels produced ranged from 0.3 to 30.0 percent of levels produced by wild-type *T. arundinaceum* (Table 5). Thus, in some species that lack other tri genes (e.g., *T. stromaticum*, *T. decipiens*, and *T. polysporum*), tri5 has retained its function in trichodiene production (Table 5). In contrast, transformants carrying tri5 homolog from *T. cf. fertile* or *T. gamsii* T065 did not produce detectable levels of HA or any of the other more than 150 trichothecene analogs detected by our analytical systems. We analyzed a total of seven transformants each with the *T. cf. fertile* and *T. gamsii* T065 homologs, and none produced detectable levels of trichothecenes (Table 5; Supplementary Figure S7). These findings indicate that in *T. cf. fertile* and *T. gamsii* T065, which lack other tri genes, the tri5 gene has not retained its function in trichodiene biosynthesis.

We confirmed expression of the tri5 homologs in transformants of Δtri5.3 using qPCR analysis with RNAs extracted from mycelia obtained from the HA production analysis. The results of the analysis indicate that the tri5 homologs from all seven donor species were expressed at high levels (Figure 8). Thus, transformants carrying the *T. cf. fertile* and *T. gamsii* T065 homologs that did not produce trichothecenes had high levels of tri5 expression.

### Production of Volatiles in Heterologous Expression Systems

To determine whether the *T. cf. fertile* or *T. gamsii* T065 tri5 homologs confer production of a terpene(s) other than trichodiene, we analyzed Δtri5.3 transformants expressing these tri5 homologs. Thus, if the *T. cf. fertile* and *T. gamsii* T065 TR15 homologs catalyze synthesis of a terpene other than trichodiene, the terpene should be detected as a volatile. However, GC–MS analysis of the headspace of cultures did not reveal any qualitative differences in volatiles produced by (i) transformants expressing the *T. cf. fertile* or *T. gamsii* T065 tri5 homolog, (ii) untransformed control strain Δtri5.3, and (iii) wild-type *T. arundinaceum*, except that the wild type produced trichodiene while the transformants and Δtri5.3 did not. These results do not provide evidence that the *T. cf. fertile* and *T. gamsii* T065 tri5 homologs confer production of trichodiene or another volatile terpene.
Analysis of Trichodiene Production by *T. decipiens*, *T. polysporum*, and *T. stromaticum*

In the heterologous expression analysis, the *T. decipiens*, *T. polysporum* and *T. stromaticum tri5* homologs complemented the *tri5* mutation in *T. arundinaceum*, indicating that the homologs are functional. To further examine their function, we determined whether wild-type strains of *T. decipiens*, *T. polysporum*, and *T. stromaticum* produced trichodiene using the headspace assay described for the previous section. No trichodiene was detected in the headspace from cultures of the three species. Thus, even though the *T. decipiens*, *T. polysporum*, and *T. stromaticum tri5* homologs are functional, activity of the corresponding enzymes could not be detected in the native strains. A likely explanation for this is the extremely low level of *tri5* expression in the three species under the growth conditions used in our experiment (Figure 6).

**Table 5** | Harzianum A (HA) production by transformants of the *T. arundinaceum tri5* deletion mutant (strain Δ*tri5.3*) expressing *tri5* homologs from other *Trichoderma* species.

| Strain          | Origin of *tri5* homolog | HA (μg/ml) | % production of positive control |
|-----------------|--------------------------|------------|----------------------------------|
| wild-type *T. arundinaceum* | *T. arundinaceum* | 168.67 ± 1.87 | 100                             |
| Δ*tri5.3*       | None                     | nd         | 0                               |
| Δ*T5_TARPUN 24* | *T. arundinaceum*        | 34.47 ± 2.12 | 20.43                           |
| Δ*T5_TARPUN 30* | *T. arundinaceum*        | 51.49 ± 0.71 | 30.52                           |
| Δ*T5_TSBAL2*    | *T. balareicum*          | 0.45 ± 0.03 | 0.27                            |
| Δ*T5_TSBAL3*    | *T. balareicum*          | 32.15 ± 6.83 | 19.06                           |
| Δ*T5_TSTDEC1*   | *T. decipiens*           | 3.12 ± 0.28 | 1.85                            |
| Δ*T5_TSTDEC3*   | *T. decipiens*           | 2.53 ± 0.22 | 1.50                            |
| Δ*T5_TTSFERT2*  | *T. cf. fertile*         | nd         | 0                               |
| Δ*T5_TTSFERT3*  | *T. cf. fertile*         | nd         | 0                               |
| Δ*T5_TSGAMS 6*  | *T. gamsii T065*         | nd         | 0                               |
| Δ*T5_TSGAMS 10* | *T. gamsii T065*         | nd         | 0                               |
| Δ*T5_TSPOLY 1*  | *T. polysporum*          | 12.62 ± 0.57 | 7.48                            |
| Δ*T5_TSPOLY 2*  | *T. polysporum*          | 11.39 ± 0.01 | 6.75                            |
| Δ*T5_TSTSTRO 4* | *T. stromaticum*         | 30.10 ± 2.3 | 17.84                           |
| Δ*T5_TSTSTRO 5* | *T. stromaticum*         | 28.08 ± 4.98 | 16.65                           |

*Percent of HA level relative to level produced by wild-type *T. arundinaceum*. 

**Table 4** | Maximum likelihood comparison between the hypotheses H0 based on a constant dN/dS ratio in the entire phylogenetic tree; and H1, with a different dN/dS ratio for the *Trichoderma* branch in the *tri5* or housekeeping gene trees.

| Strain | Sample | Expression ratio | p(H1) |
|--------|--------|------------------|-------|
| 1. *T. arundinaceum* | 14.47* | 0.000            |
| 2. *T. balareicum* | 0.007* | 0.000            |
| 3. *T. cf. fertile* | -       | -                |
| 4. *T. decipiens* | 0.010* | 0.000            |
| 5. *T. stromaticum* | 0.016* | 0.000            |
| 6. *T. polysporum* | 0.021* | 0.000            |
| 7. *T. gamsii T065* | 0.004* | 0.000            |

**Figure 6** | *tri5* expression in selected *Trichoderma* species. The data are derived from Real Time quantitative qPCR analysis and values are the level of expression of *tri5* versus the actin gene. Statistically significant expression ratio values [p(H1)<0.05] are indicated with an asterisk. nd indicates not detected.

The statistical differences between the likelihood values observed for each species pair was analyzed by the Kishino-Hasegawa (pKH) test, using the program CodeML in the PAML software package.

*The value in bold text is the highest ω value observed in the *Trichoderma* branch (ω₉) versus in all branches (ω₀) in the *tri5* tree.*

*Indicates p values of 0.011-0.049; and **indicate p values < 0.01.*
Analysis of the Antifungal Activity
Production of trichothecenes has been shown to contribute to the ability of *T. arundinaceum* and *T. brevicompactum* to inhibit the growth of some plant pathogenic fungi. Therefore, we examined the ability of other trichothecene-producing and nonproducing species/strains of *Trichoderma* to inhibit the growth of the plant pathogen *Rhizoctonia solani*. In all experiments, the *T. arundinaceum* Δtri5 mutant, strain Δtri5.3, caused markedly less growth inhibition than its wild-type progenitor strain, indicating that trichothecene production contributes substantially to the antifungal activity of *T. arundinaceum* against *R. solani* (Supplementary Figures S8, S9).

In one set of experiments, we assessed antifungal activity of wild-type strains of four species that have tri5 but no other tri genes: *T. decipiens*, *T. gamsii* T065, *T. polysporum*, and *T. stromaticum*. All four species inhibit *R. solani* growth much less than wild-type *T. arundinaceum* (Supplementary Figure S8). Growth inhibition caused by *T. decipiens* was similar to that caused by Δtri5.3, while inhibition caused by *T. gamsii* T065, *T. polysporum*, and *T. stromaticum* was less than that caused by Δtri5.3.

In a second set of experiments, we examined the effect that expression of *tri5* homologs from trichothecene-producing and nonproducing species of *Trichoderma* had on the antifungal activity of strain Δtri5.3. The antifungal activity of Δtri5.3 transformants expressing *tri5* homologs was correlated with the levels of HA produced by them (Supplementary Figure S9). That is, transformants expressing the *T. arundinaceum* (positive control), *T. balearicum*, and *T. stromaticum tri5* homologs, which produced relatively high levels of trichothecenes, caused the highest levels of inhibition, while transformants expressing the *T. decipiens* and *T. polysporum tri5* homologs, which produce lower levels of HA, cause less growth inhibition (Supplementary Figure S9). Further, transformants expressing *tri5* homologs from *T. cf. fertile* and *T. gamsii* T065, which did not produce HA, caused levels of inhibition similar to those caused by strain Δtri5.3 (Supplementary Figure S9).
DISCUSSION

Distribution and Location of tri5 in Trichoderma Lineages

Various combinations of tri genes sufficient to confer trichothecene biosynthesis have been reported in 14 fungal genera of the Ascomycota (Proctor et al., 2020), and trichothecene production has been confirmed in species from 10 of these genera, including at least six species of Trichoderma (Cardoza et al., 2011; Tijerino et al., 2011; Dawidziuk et al., 2014; Sun et al., 2016; Chen et al., 2018). In the current study, examination of genome sequences of 35 Trichoderma species revealed that tri5 and the tri cluster occur in diverse Trichoderma species but that tri5 is more common than the tri cluster (Table 1).

![FIGURE 8 | tri5 expression in T. arundinaceum strains generated in the complementation analysis. Values are derived from Real Time quantitative PCR (qPCR) analysis and are the ratio of the level of expression of tri5 versus the actin gene. Strains: wild type – wild-type T. arundinaceum, and Δtri5.3 (Δ5) – T. arundinaceum Δtri5 deletion mutant. The 14 other strains are transformants of Δtri5.3 expressing tri5 homologs from seven Trichoderma species. Statistically significant expression ratio values \( p(H1) < 0.05 \) are indicated with an asterisk. nd indicates not detected.](image-url)
The results also indicated that physical separation of tri5 and the tri cluster is a conserved character among Trichoderma species that have both loci. It is noteworthy that in 18 Trichoderma species, tri5 was located at the same genomic location (GR1) as originally described in T. arundinaceum and T. brevicipactum, whereas the occurrence of tri5 in other genomic location was much less common. Nevertheless, its occurrence at GR2–GR5 suggests that tri5 has undergone two translocation events during divergence of Trichoderma species. However, our understanding of GR2–GR5 was limited by the small contigs on which these regions occurred. Although our analyses demonstrated conservation of the physical separation of tri5 and the tri cluster in Trichoderma, the biological significance of the separation is not clear (Proctor et al., 2020).

The 35 Trichoderma genome sequences examined in the current study represent 12 phylogenetically distinct lineages of the genus Trichoderma, and the species tree inferred in the current study (Figure 1) was correlated with previously reported species trees (e.g., Kubicek et al., 2019). Analysis of the presence and absence of tri5 in the context of the species tree revealed that the gene was present in all lineages examined except Deliquescens and Longibrachiatum (Table 1). Within the 10 lineages with tri5, its occurrence varied from present in all species examined (e.g., Psychrophila), present in most species (Brevicompactum), and present in a minority of species (Harzianum/Virens). This variation in occurrence suggests that tri5 has been lost multiple times in Trichoderma and potentially multiple times within some lineages. However, the results do not rule out the possibility that horizontal transfer between Trichoderma species has contributed to variation in occurrence of tri5 within and among lineages. Regardless, its widespread distribution among the lineages examined suggests that tri5 was present in Trichoderma prior to divergence of the lineages.

Fusarium trichothecenes are an important agricultural and food/feed safety concern because of their toxicity and frequent occurrence in crops (Munkvold et al., 2021). A comparison of results from the current study on Trichoderma and previous studies on Fusarium indicate marked difference in the presence and absence of tri genes and trichothecene production in the two genera. When the genus Trichoderma is viewed as a whole, the distribution of tri5, the tri cluster, and trichothecene production are discontinuous. In Fusarium by contrast, the presence of the tri cluster, which includes tri5, and trichothecene production is continuous in two closely related, multispecies lineages (the F. incarnatum-equiseti and F. sambucinum species complexes), whereas the cluster and production are absent in 20 other lineages (Proctor et al., 2009; O'Donnell et al., 2013; Villani et al., 2019; Brown et al., 2020; Laraba et al., 2021).

A potential caveat is that Fusarium may encompass more phylogenetic diversity than Trichoderma (Crous et al., 2021; Geiser et al., 2021). Nevertheless, the F. incarnatum-equiseti and F. sambucinum species complexes comprise at least 35 and 70 phylogenetically distinct species, respectively, in which the tri cluster appears to be continuously distributed (Xia et al., 2019; Laraba et al., 2021). Based on our current understanding, tri5 and the tri cluster do not have a continuous distribution over similar numbers of phylogenetically distinct Trichoderma species. This difference in distribution suggests differences in selection to retain tri genes in the two genera. What the drivers of such differences in selection would be are not clear, but one tempting argument is that Fusarium species tend to be plant pathogens while Trichoderma species tend to be saprobes. This argument falls short, however, because some members of the F. incarnatum-equiseti and F. sambucinum complexes are weak plant pathogens at best, and some other Fusarium species complexes include species that are highly aggressive plant pathogens but lack tri genes (Leslie and Summerell, 2006; Munkvold et al., 2021).

Evidence for a Positive Selection of tri5 in Trichoderma spp.

The results of the current study also provide evidence for positive selection of the ancestral Trichoderma tri5 during its divergence from tri5 in other fungi (Figures 4, 5). There is evidence for positive selection of other secondary metabolite biosynthetic genes in fungi (Massonnet et al., 2018; Alouane et al., 2021). In these other fungi, the selection was attributed to adaptation to environmental changes. Such adaptation could have also driven positive selection of tri5, but it is not clear what the specific driver(s) would be. Three other observations could provide further insight into the positive selection and other aspects of the evolution of Trichoderma tri5. First, as noted previously, tri5 and the tri cluster are at different genomic locations in Trichoderma. Further, there is no evidence for positive selection of Trichoderma tri cluster genes; the Trichoderma branch in trees inferred from each of six tri cluster genes is markedly shorter than in the tri5 tree (Proctor et al., 2018). Their physical separation and different rates of divergence suggest that the Trichoderma tri5 and tri cluster have had distinct evolutionary histories. Second, in the tri5 tree, branch lengths within three single-genus clades, Trichoderma, Fusarium, and Stachybotrys, are generally similar, although there are some differences (Figure 4). This similarity suggests that the positive selection occurred on the ancestral Tri. tri5 but was relaxed as Trichoderma species diverged from one another. Third, we surmise that the location of tri5 at GR1 is ancestral to its location at GR2–GR5, because tri5 occurs at GR1 in seven diverse Trichoderma lineages, while tri5 occurs in GR2–GR5 in only three lineages, two of which are closely related (Figure 2). Further, despite evidence that positive selection on a gene can affect divergence of neighboring genes (Barton, 2000), we found no evidence for positive selection of three tri5-flanking genes at GR1 (Supplementary Figure S6). A possible explanation for this is that the positive selection on tri5 occurred before it was located at GR1.

The above observations suggest that the evolutionary history of the Trichoderma tri5 has been complex. Nevertheless, we propose a relatively simple three-step evolutionary model that accounts for the observations. Step 1 – a common ancestor of extant Trichoderma lineages had a tri cluster that included tri5, but subsequently lost tri5 through gene deletion or pseudogenization. Step 2 – the Trichoderma ancestor reacquired
tris5 via horizontal transfer from a fungus that occupied a markedly different habitat than other trichothecene-producing fungi for some of its evolutionary history. This is consistent with evidence for positive selection on tris5. Step 3 – when it was horizontally transferred, tris5 integrated at GR1 in the ancestral Trichoderma genome. This model accounts for the observations noted above, and we have found precedence for each step of the model. For example, the absence of tris4 in Spicellum roseum provides precedence for loss (Step 1) and subsequent reacquisition (Step 2) of an early biosynthetic step essential for formation of trichothecenes \( (Proctor \text{ et al., } 2018). \) Further, there is an increasing body of evidence for horizontal transfer of secondary metabolite biosynthetic genes among fungi (Step 2; Slot and Rokas, 2011; Proctor et al., 2013; Villani et al., 2019). Although there is precedence for positive selection and horizontal transfer of secondary metabolite biosynthetic genes in fungi, we are not aware of an example of transfer of a gene that has undergone positive selection prior to the transfer. In 2014, there were publications on genome sequences for trichothecene-producing species of only two fungal genera: Fusarium and Stachybotrys \( (Ma \text{ et al., } 2010; \text{ Semeiks et al., } 2014). \) Today, there are publications on genome sequences of trichothecene-producing species in at least 12 genera \( (Pitt \text{ et al., } 2017; \text{ Proctor et al., } 2020; \text{ Geiser et al., } 2021). \) As more genome sequences are generated and analyzed, they are likely to provide further insight into the evolutionary history of tris5 and other tri genes in Trichoderma, and facilitate refinement or rejection of the model proposed above.

**Genomic Location and Functionality of Trichoderma tris5**

Although tris5 occurred at different genomic locations in some Trichoderma species, it was at the same location (GR1) in all species that had the tri cluster and that produced trichothecenes. The four species that had tris5 at other genomic locations (GR2–GR5), lacked the tri cluster and, therefore, did not produce trichothecenes (Figure 2; Table 1). Whether this correlation is biologically significant is not known, but the correlation raises the question, is the presence of tris5 at location GR1 more favorable for trichothecene production than its presence at other genomic locations? If the answer is yes, the occurrence of tris5 at locations GR2–GR5 could represent a relaxation of selection for its occurrence at GR1. The T. gamsii T065 and T. polysporum tris5 homologs, which occur at GR3 and GR4 respectively, differed in their ability to complement the T. arundinaceum \( \Delta \)tris5 mutant. Thus, the occurrence of tris5 at locations other than GR1 has not necessarily caused tris5 to lose its function in trichodiene biosynthesis.

Of the nine Trichoderma species that have tris5 but lack the tri cluster, T. cf. fertile, T. margarectense, and T. rhododendri have tris5 homologs with 1–3 point mutations that introduce internal stop codons or a frameshift and, as a result, have potential to render the TRIS5 protein nonfunctional (Supplementary Figure S10). By contrast, six of the nine species have tris5 homologs that are apparently functional based on their DNA sequences, and further the function of the T. decipiens, T. polysporum, and T. stromaticum tris5 homologs was demonstrated by their ability to complement the T. arundinaceum \( \Delta \)tris5 mutant. The presence of a functional tris5 homolog in some Trichoderma species that lack the tri cluster raises the question, what is the function of tris5 in the absence of trichothecene production? Our analyses of volatiles produced by transformants of the T. arundinaceum \( \Delta \)tris5 mutant carrying various tris5 homologs failed to provide support for the hypothesis that the homologs confer the ability to produce a terpene(s) other than trichodiene. A possible alternative explanation is evident from a study in which the T. arundinaceum tris5 was overexpressed in the biocontrol agent \( T. \) harzianum, which normally lacks tris5 and the tri cluster \( (Table \ 1; \text{ Taylor et al., } 2021). \) The tris5-expressing \( T. \) harzianum strain produced trichodiene and had enhanced biocontrol activity against Fusarium graminearum, a trichothecene-producing fungus that is a predominant cause of ear rot of maize and Fusarium head blight of wheat and barley. In addition, tris5-expressing \( T. \) harzianum volatiles, which included trichodiene, downregulated tri gene expression and trichothecene production in \( F. \) graminearum \( (Taylor \text{ et al., } 2021). \) Thus, we propose the following model: in Trichoderma species that have a functional tris5 but lack the tri cluster, trichodiene production provides a competitive advantage by inhibiting production of trichothecenes and/or other toxic terpenoids in other fungi.

**Trichothecene Biosynthesis in Trichoderma Species**

Based on knowledge of the roles of the various tri genes in trichothecene biosynthesis, we have proposed a biosynthetic scheme to explain the biochemical relationships of trichothecene analogs that have been reported to be produced by Trichoderma species \( (Figure \ 9). \) Although trichodermin is a pathway endpoint in some species such as T. brevicipactum \( (Tijerino \text{ et al., } 2011), \) in the proposed scheme, trichodermin is also an intermediate from which four pathway branches arise. That trichodermin is an intermediate in HA biosynthesis has been demonstrated previously \( (Lindo \text{ et al., } 2019). \) Given the similarity in structure the octa-2,4,6-trienedioyl substituent of HA to the 8-carbon polyketide portion of the macroclide ring of roridins, it is possible that trichodermin is also an intermediate in roridin biosynthesis as well. With the exception of the tris17-encoded polyketide synthase, the genes that confer macroclide ring formation in Trichoderma and other fungi are not known \( (Semeiks \text{ et al., } 2014; \text{ Proctor et al., } 2018; \text{ Zhu et al., } 2020). \) Formation of 4-acetyltrichothecolone and 16-hydroxytrichodermin from trichodermin would require only one or two biochemical reactions \( (Figure \ 9). \) In Fusarium species, oxygenation of trichothecenes at carbon atom 8 \( (C8) \) is catalyzed by a tris1-encoded cytochrome P450 monooxygenase. Analysis of the T. psychrophilum genome sequence did not reveal the presence of a tris1 homolog. Therefore, a gene other than tris1 must confer 8-oxygenation in T. psychrophilum. As far as we are aware, a tri gene that confers 16-hydroxylation has not been identified in any trichothecene-producing fungus. Thus, the gene required for formation of 16-hydroxy substituent of 16-hydroxytrichodermin produced by T. albolutelescens remains to be determined \( (Ryu \text{ et al., } 2017). \)
In the biosynthetic scheme, we proposed that the formation of isotrichodermin occurs via 3-hydroxylation of trichodiene rather than 3-hydroxylation of EPT because the latter modification occurs via the activity of TRI4 during trichothecene biosynthesis in Fusarium species. That is, Fusarium TRI4 homologs catalyze hydroxylation of trichodiene at C2, C3, C11, and C13, which causes a nonenzymatic cyclization resulting in the formation of isotrithodermol (3-hydroxy EPT; McCormick et al., 2006; Tokai et al., 2007). The tri101-encoded acetyltransferase then catalyzes acetylation of the 3-hydroxy substituent to form isotrichodermin (3-acetyl EPT; McCormick et al., 1999). The activity of TRI4 marks a fundamental difference in trichothecene biosynthesis in Trichoderma and Fusarium. Analysis of T. arundinaceum tri4 indicates that the Trichoderma TRI4 catalyzes hydroxylation of trichodiene at only C2, C11, and C13, resulting in 6,7-dihydroxy-2,6-octadienoyl-CoA (Cardoza et al., 2015). This activity is consistent with the absence of a 3-oxygen atom in all reported Trichoderma trichothecenes except for isotrichodermin produced by T. albolutescens. It is possible, however, that the T. albolutescens TRI4 homolog has low levels of trichodiene 3-hydroxylation activity, as is the case for the Myrothecium roridum TRI4 (McCormick and Alexander, 2007). Further, because many fungi have trichothecene 3-acetylation activity (Khatibi et al., 2011), it would not be unusual for T. albolutescens to also have such activity. Thus, we propose that production of isotrichodermin (3-acetyl EPT) by T. albolutescens results from 3-hydroxylation activity of its TRI4 homolog and a 3-acetylation activity that is relatively common among fungi.

**Conclusion**

The results of this study expand the understanding of trichothecene production in filamentous fungi by demonstrating that production is discontinuously distributed over a wide breadth of phylogenetic diversity of the genus Trichoderma. This distribution of trichothecene production among Trichoderma species suggests the toxins could play a role in the ability of

![Proposed biosynthetic pathway for Trichoderma trichothecenes.](https://example.com/pathway.png)

**FIGURE 9** | Proposed biosynthetic pathway for Trichoderma trichothecenes. Arrows correspond to biosynthetic steps and enzyme activities already described, except for those labeled with question marks, which point out the genes not yet identified or characterized. Chemical structures shaded in yellow correspond to the final trichothecene analogs detected in the Trichoderma species analyzed in the present work and included in Table 1. Genes are indicated in bold and black lowercase letters in italics, and enzymes are indicated by bold and blue uppercase letters.
some species to survive and compete under certain environmental conditions. The finding that functional homologs of tri5 are retained in some species suggests that trichodiene production in the absence of trichothecene production could also provide an ecological advantage. This is turn consistent with the hypothesis proposed by Taylor et al. (2021) that trichodiene can act as a signaling molecule. Further, the genome sequences of Trichoderma species that produce unusual trichothecene analogs, such as 4-acetyltrichothecolone and 16-hydroxytrichodermcin, should provide a source to identify novel trichothecene biosynthetic genes that could provide insight into the ecological roles of trichothecenes in this important genus of microorganisms.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

AUTHOR CONTRIBUTIONS

SG and RP conceived the work, designed the experiments, performed genome sequencing, participated in bioinformatics analyses, and coordinated the work of the rest of the coauthors. SM, RC, H-SK, MB, LY, MV, PC, and GC-H performed the chemical studies, fungal transformations, tri5 mutant isolation and complementation, and antifungal experiments, and participated in the bioinformatics analyses. WJ provided fungal material. W-YZ and CW participated in the fungal genome sequencing. All authors collaborator in the discussion of the results, and contributed to the writing and to the corrections made to reach the final version of this manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online https://www.frontiersin.org/articles/10.3389/fmicb.2021.791641/full#supplementary-material.

REFERENCES

Alouane, T., Rimbert, H., Bornmann, J., González-Montiel, G. A., Loesgen, S., Schäfer, W., et al. (2021). Comparative genomics of eight Fusarium graminearum strains with contrasting aggressiveness reveals an expanded open pan genome and extended effector content signatures. Int. J. Mol. Sci. 22:6257. doi: 10.3390/ijms22126257
Altschul, S. F., Gish, W., Miller, W., Myers, E. W., and Lipman, D. J. (1990). Basic local alignment search tool. J. Mol. Biol. 215, 403–410. doi: 10.1016/S0022-2836(05)80360-2
Bankevich, A., Nurk, S., Antipov, D., Gurevich, A. A., Dvorkin, M., Kulikov, A. S., Lesin, V. M., Nikolenko, I. I., Pham, S., Proužek, A., et al. (2012). SPAdes: a new genome assembly algorithm and its applications to single-cell sequencing. J. Comput. Biol. 19, 455–477. doi: 10.1089/cnb.2012.0021
Barton, N. H. (2000). Genetic hitchhiking. Philos. Trans. R. Soc. Lond. B 355, 1553–1562. doi: 10.1098/rstb.2000.0716
Blin, K., Shaw, S., Kloosterman, A. M., Charlop-Powers, Z., van Wezel, G. P., Medema, M. H., et al. (2021). antiSMASH 6.0: improving cluster detection and comparison capabilities. Nucleic Acids Res. 49, W29–W35. doi: 10.1093/nar/gkab335
Brown, D. W., Villani, A., Susca, A., Moretti, A., Hao, G., Kim, H. S., et al. (2020). Gain and loss of a transcription factor that regulates late trichothecene production in the plant defense response and fungal physiology: overexpression of the Trichoderma arundinaceum tri4 gene in T. harzianum. Appl. Environ. Microbiol. 81, 6355–6366. doi: 10.1128/AEM.01626-15
Cardoza, R. E., McCormick, S. P., Lindo, L., Kim, H.-S., Olivera, E. R., Nelson, D. R., et al. (2019). A cytochrome P450 monoxygenase gene required for biosynthesis of the trichothecane toxin harzianum A in Trichoderma. Appl. Microbiol. Biotechnol. 103, 8087–8103. doi: 10.1007/s00253-019-10047-2
Cardoza, R. E., McCormick, S. P., Malmierca, M. G., Olivera, E. R., Alexander, N. J., Monte, E., et al. (2015). Effects of trichothecene production on the plant defense response and fungal physiology: overexpression of the Trichoderma arundinaceum tri4 gene in T. harzianum. Appl. Environ. Microbiol. 81, 6355–6366. doi: 10.1128/AEM.01626-15
Cardoza, R. E., Vizcaino, J. A., Hermosa, M. R., Monte, E., and Gutiérrez, S. (2006). A comparison of the phenotypic and genetic stability of recombinant Trichoderma spp. generated by protoplast- and agrobacterium-mediated transformation. J. Microbiol. 44, 383–395.
Casqueiro, J., Gutiérrez, S., Batuelos, O., Hijarrubia, M. J., and Martín, J. F. (1999). Gene targeting in Penicillium chrysogenum: disruption of the lys2 gene leads to penicillin overproduction. J. Bacteriol. 182, 1181–1188. doi: 10.1128/JB.181.4.1181-1188.1999
Chen, B., Li, E., Liu, L., Liao, M., Zhu, Z., Zhuang, W., et al. (2018). Botryane sesquiterpenoids, cyclopentadepsipeptides, xhanthones, and trichothecenes from Trichoderma oligospororum. Planta Med. 84, 1055–1063. doi: 10.1055/a-0593-6603
Cole, R. J., Dorner, J. W., Cox, R. H., Cunfer, B. M., Cutler, H. G., and Stuart, B. P. (1981). The isolation and identification of several trichothecene mycotoxins from Fusarium heterosporum. J. Nat. Prod. 44, 324–330. doi: 10.1021/np50015a015
Crous, P. W., Lombard, L., Sandoval-Denis, M., Seifert, K. A., Schroers, H. J., Chaverri, P., et al. (2021). Fusarium: more than a node or a foot-shaped basal cell. Stud. Mycol. 98, 100116–100116. doi: 10.1016/j.studmyco.2021.100116
relative expression results in real-time PCR. *Nucleic Acids Res.* 30:x36. doi: 10.1093/nar/30.3.x36

Pitt, J. I., Lange, L., Lacey, A. E., Vuong, D., Midgley, D. J., Greenfield, P., et al. (2017). *Aspergillus hancockii* sp. nov., a biosynthetically talented fungus endemic to southeastern Australian soils. *PloS One* 12:e0170254. doi: 10.1371/journal.pone.0170254

Proctor, R. H., Desjardins, A. E., Plattner, R. D., and Hohn, T. M. (1999). A polyketide synthase gene required for biosynthesis of fumonisin mycotoxins in *Gibberella fujikuroi* mating population A. *Fungal Genet. Biol.* 27, 100–112. doi: 10.1006/fgbi.1999.1141

Proctor, R. H., McCormick, S. P., Alexander, N. J., and Desjardins, A. E. (2009). Evidence that a secondary metabolic biosynthetic gene cluster has been grown by gene relocation during evolution of the filamentous fungus *Fusarium*. *Mol. Microbiol.* 74, 1128–1142. doi: 10.1111/j.1365-2958.2009.06927.x

Proctor, R. H., McCormick, S. P., and Gutiérrez, S. (2020). Genetic bases for variation in structure and biological activity of trichothecene toxins produced by diverse fungi. *Appl. Microbiol. Biotechnol.* 104, 5185–5199. doi: 10.1007/s00253-020-10616-2

Szwalbe, A. J., Williams, K., Song, Z., de Mattos-Shipley, K., Vincent, J. L., Bailey, A. M., et al. (2019). Characterisation of the biosynthetic pathway to agestins A and B reveals the reductive route to chrysophanol in fungi. *Chem. Sci.* 10, 233–238. doi: 10.1039/C8SC03776G

Taylor, L., Gutiérrez, S., McCormick, S. P., Bakker, M. G., Proctor, R. H., Teresi, J., et al. (2021). Use of the volatile trichodiene to reduce Fusarium head blight and trichothecene contamination in wheat. *Microb. Biotechnol.* doi: 10.1111/1751-7915.13742

Tijerino, A., Cardoza, R. E., Moraga, J., Malmierca, M. G., Vicente, E., Aleu, J., et al. (2011). Overexpression of the trichodiene synthase gene *tri5* increases trichodermin production and antimicrobial activity in *Trichoderma brevicompactum*. *Fungal Genet. Biol.* 48, 285–296. doi: 10.1016/j.fgb.2010.11.012

Tokai, T., Koshino, H., Takahashi-Ando, N., Sato, M., Fujimura, M., and Kimura, M. (2007). *Fusarium Tri4* encodes a key multifunctional cytochrome P450 monooxygenase for four consecutive oxygenation steps in trichothecene biosynthesis. *Biochem. Biophys. Res. Commun.* 353, 412–417. doi: 10.1016/j.brc.2006.12.033

Vaidya, G., Lohman, D. J., and Meier, R. (2011). SequenceMatrix: concatenation software for the fast assembly of multi-locus datasets with character set and codon information. *Cladistics* 27, 171–180. doi: 10.1111/j.1096-0031.2010.00329.x

Villani, A., Proctor, R. H., Kim, H. S., Brown, D. W., Logrieco, A. F., Amatulli, M. T., et al. (2019). Variation in secondary metabolite production potential in the *Fusarium incarnatum-epsatus* species complex revealed by comparative analysis of 13 genomes. *BMC Genomics* 20:314. doi: 10.1186/s12864-019-5567-7

Vincente, I., Baronecelli, R., Morán-Diez, M. E., Bernardi, R., Puntoni, G., Hermosa, R., et al. (2020). Combined comparative genomics and gene expression analyses provide insights into the terpene synthases inventory of *Trichoderma*. *Microorganisms* 8:603. doi: 10.3390/microorganisms810603

Xia, J. W., Sandoval-Denis, M., Crous, P. W., Zhang, X. G., and Lombard, L. (2019). Numbers to names – restyling the *Fusarium incarnatum-epsatus* species complex. *Persoonia* 43, 186–221. doi: 10.3767/persoonia.2019.43.05

Yang, Z. (2007). PAML 4: a program package for phylogenetic analysis by maximum likelihood. *Mol. Biol. Evol.* 24, 1588–1591. doi: 10.1093/molbev/msm088

Zhang, J., Rosenberg, H. F., and Nei, M. (1998). Positive Darwinian selection alter gene duplication in primate ribonuclease genes. *Proc. Natl. Acad. Sci. U. S. A.* 95, 3708–3713. doi: 10.1073/pnas.95.7.3708

Zhu, M., Cen, Y., Ye, W., Li, S., and Zhang, W. (2020). Recent advances on macrocyclic trichotheccenes, their bioactivities and biosynthetic pathway. *Toxins* 12:417. doi: 10.3390/toxins12060417

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