Identification of a Novel Partitivirus of *Trichoderma harzianum* NFCF319 and Evidence for the Related Antifungal Activity

Jeesun Chun¹, Han-Eul Yang² and Dae-Hyuk Kim¹,²*

¹ Institute for Molecular Biology and Genetics, Chonbuk National University, Jeonju, South Korea, ² Department of Bioactive Material Sciences, Chonbuk National University, Jeonju, South Korea

We have reported 15 agarose gel band patterns of double-stranded RNA (dsRNA) from *Trichoderma* spp. We describe herein that band pattern IX in *Trichoderma harzianum* NFCF319, which appeared to be a single band but consisted of two dsRNAs of similar size, was identified as a novel mycovirus, designated *Trichoderma harzianum* partitivirus 1 (ThPV1). The larger segment (dsRNA1) of the ThPV1 genome comprised 2,289 bp and contained a single open reading frame (ORF) encoding an RNA-dependent RNA polymerase (RdRp). The smaller segment (dsRNA2) consisted of 2,245 bp with a single ORF encoding a capsid protein (CP). Evaluation of the deduced amino acid sequence and phylogenetic analysis indicated that ThPV1 is a new member of the genus *Betapartitivirus* in the family *Partitiviridae*. Curing of virus infection by single-sporing generated 31 virus-free single-spore clones. No significant differences in growth rate, conidia production, or pigmentation were observed between ThPV1-infected and -cured isogenic strains. In addition, comparison of the newly ThPV1-transmitted isolates with their ThPV1-cured parental strain showed no significant difference in colony morphology or pigmentation. However, inhibition of growth in co-cultured *Pleurotus ostreatus* and *Rhizoctonia solani* by *T. harzianum* was increased in ThPV1-containing strains compared with ThPV1-cured isogenic strains. Moreover, β-1,3-glucanase activity was significantly increased in the ThPV1-containing strains. However, no difference in chitinase activity was observed, suggesting that ThPV1 regulates the activity of a specific fungal enzyme.

**Keywords:** *Trichoderma harzianum*, mycovirus, partitivirus, mycoparasitism, antifungal activity

**INTRODUCTION**

Mycoviruses, i.e., fungal viruses, are widespread and have been found in all major taxa of filamentous fungi and yeasts (Van Alfen, 1986; Nuss and Koltin, 1990; Wickner, 1992). The majority of characterized mycoviruses have a double-stranded RNA (dsRNA) genome, although others have single-stranded RNA (ssRNA) and DNA genomes. Mycoviruses of seven families, *Chrysoviridae*, *Endornaviridae*, *Megabirnaviridae*, *Quadirviridae*, *Partitiviridae*, *Reoviridae*, and *Totiviridae* have dsRNA genomes, while those of six families, *Alphaflexiviridae*, *Barnaviridae*, *Gammaflexiviridae*, *Hypoviridae*, *Narnaviridae*, and *Mymonaviridae* have ssRNA genomes according to a report by the International Committee on the Taxonomy of Viruses (ICTV) in 2016 (Ghabrial et al., 2015).
Using multi-omics techniques, a large number of novel mycoviruses are expected to be discovered and characterized from various fungi. Recently, we reported the presence of a variety of dsRNA elements in many strains of Trichoderma spp. causing a green mold disease in shiitake mushroom (Lentinula edodes) and confirmed the presence of novel mycoviruses in these Trichoderma spp. (Yun et al., 2016). Although infection of fungi by many mycoviruses is asymptomatic or cryptic (Ghabrial, 1998), there are many cases of mycoviruses inducing viral-specific symptoms in the host and reduced fungal virulence, known as hypovirulence, is one of representative examples in phytopathogenic fungi (Nuss, 2005).

Members of the genus Trichoderma are fast-growing ubiquitous soil fungi. The evident success of Trichoderma species is due to the characteristic mechanisms of Trichoderma to survive and proliferate, which include an aggressive ability to inhibit other fungi and production of various enzymes for the degradation of complex carbohydrates (Samuels and Hebar, 2015). These activities have been exploited for human benefit; indeed, some Trichoderma species are used as biocontrol agents, and in the production of commercial enzymes and secondary metabolites, and are endosymbionts for their host plant (Kubicek and Penttila, 1998; Sivasithamparam and Ghisalberti, 1998; Hatvani et al., 2002; Howell, 2003). T. harzianum is probably the most commonly cited species and is widely recognized as a potential biocontrol component in controlling common soil-borne plant pathogens (i.e., Fusarium, Pythium, and Rhizoctonia), but is also reported as the causal agent of green mold disease in cultivated mushrooms (Papavizas, 1985; Tokimoto, 1985; Ulhoa and Peberdy, 1992; Seaby, 1998; Kredics et al., 2010). Thus, changes in the beneficial or pathological characteristics of T. harzianum impact the industrial application or prevention of this fungus. We have reported the presence of dsRNA in T. harzianum; however, the nature and biological function of the dsRNA are unclear.

In this study, we characterized the dsRNA of a novel mycovirus and assessed its biological effects on T. harzianum.

**MATERIALS AND METHODS**

**Fungal Strains and Growth Condition**

Fungal strains were maintained at 25°C in the dark on PDA. Virus-containing and -cured T. harzianum strains were maintained by the hyphal-tipping technique.

**Nucleic Acid Extraction and Viral Genome Sequencing**

dsRNA extraction and Northern hybridization analysis were performed as described by Park et al. (2008). Purified dsRNA was subjected to cDNA library construction and genome sequencing on the Illumina Hiseq 2000 platform (Macrogen Inc., Seoul, South Korea). The Illumina adapter sequence reads were quality checked by FastQC and trimmed by Trimmomatic (ver. 0.32). Qualified reads were assembled to generate contigs by Trinity and abundance was estimated using RSEM software to calculate the FPKM-values.

**Rapid Amplification of cDNA Ends (RACE) Analysis**

RNA ligase-mediated rapid amplification of cDNA ends (RLM-RACE) was performed to determine the 5′- and 3′-terminal sequences of dsRNA using an RLM-RACE kit (Ambion, Austin, TX, United States). Purified dsRNA was denatured in dimethylsulfoxide and treated with calf intestine alkaline phosphatase (CIP) and tobacco acid pyrophosphatase (TAP) to remove free 5′ phosphates and cap structures. The 5′ RNA adapter oligonucleotide (5′-GCUGAUGGCGAUGAAACACUGCUUGGCUU UGAUGAAA-3′) was ligated to the decapped RNA using T4 RNA ligase. The ligates were subjected to random-primed reverse transcription and the 5′ end of a specific sequence was amplified. The 3′ terminus of dsRNA was ligated to a 3′ RACE adapter oligonucleotide (5′-GCGAGCACAGATTATACGACTCAGTATAGT12VN-3′) and subjected to RT-PCR. The resulting cDNA was amplified by PCR to determine the 3′ end sequence.

**Sequence Analysis**

Phylogenetic trees were constructed by the maximum-likelihood methods (Fitch, 1971) with the software package MEGA7 (Kumar et al., 2016) after performing multiple sequence alignments using CLUSTAL X (ver. 2.1) (Thompson et al., 1997).

**Purification of Virus Particles and Transmission Electron Microscopy**

Trichoderma harzianum betapartitivirus 1 (ThPV1) particles were purified by the method of Wang et al. (2014) and subjected to sucrose gradient ultracentrifugation. The structure of the virus-like particles was visualized by TEM on an H-7650 instrument installed in the Center for University-Wide Research Facilities (CURF) at Chonbuk National University (Hitachi, Tokyo, Japan) after negative staining with 2% uranyl acetate.

**Antifungal Activity Assay**

Conidial suspensions (1 × 10⁵ spores/ml) of T. harzianum NFCF319 were inoculated in 150 ml of potato dextrose broth (PDB) and incubated in shaking flasks for 2 days at 25°C and 180 rpm. The culture medium was passed through a 0.2 μm membrane filter and tested for each filtrate diluted with PDB containing 10⁻¹, 10⁻², and 10⁻³ of the original filtrate. Pleurotus ostreatus (ASI No. 2792) and Rhizoctonia solani AG-1 (KACC40101) were cultured by placing mycelial plugs in the center of PDA containing each filtrate, and colonial growth was assessed as the mean radial area.

**Chitinase and β-1,3-Glucanase Assay**

Culture supernatants were harvested and subjected to chitinase assays according to the manufacturer’s instructions (Sigma, St. Louis, MO, United States). β-1,3-glucanase assays were
performed in 0.05 M sodium citrate buffer (pH 4.5) with β-1,3-glucan for 2 h. The reaction was stopped by heating at 100°C for 5 min, and the amount of reducing sugar liberated was measured using neocuproine. One unit (U) of β-1,3-glucanase activity was defined as the amount of an enzyme that produced 1 µmol of reducing sugar per minute under the assay conditions.

**Transformation**

Protoplasts were prepared as described previously (Penttilä et al., 1987). Transformation using protoplasts was performed as described previously, with slight modification (Churchill et al., 1990; Kim et al., 2002). Briefly, polyethylene glycol (PEG) 6,000 (Sigma) was used instead of PEG 4,000 when transforming DNA was mixed with protoplasts. The protoplasts were transformed with pDH25, which carried the hygromycin phosphotransferase gene cassette (hph) (Cullen et al., 1987). Transformants were selected from agar plates that were supplemented with 150 µg/ml hygromycin B (Calbiochem; Merck, Darmstadt, Germany), passaged three or four times on selective medium, and single-spore isolated as described previously (Kim et al., 2002). PCR and Southern blot analysis were conducted with genomic DNA from the transformants to confirm the hygromycin B resistance gene.

**Transmission of dsRNA Virus**

Virus transmission was performed as described previously (Kim et al., 2013). Briefly, mycelial plugs of the virus-containing strain were placed on PDA medium adjacent to mycelial plugs of the virus-free hygromycin-B-resistant recipient. After 3 days of co-culture, putatively fused mycelia at the recipient border between each pair of strains were transferred to hygromycin-containing PDA, successively transferred to fresh hygromycin-containing
medium at least three times, and examined for the presence of virus. Then, the virus-containing mycelial progenies were single-spored to select for the virus-infected recipient transformants. The presence of mycovirus was confirmed by purification of dsRNA from the single-spored isolates.

RESULTS
Sequence Determination and Genome Organization
The dsRNA extracted from the mycelia of *T. harzianum* NFCF319 was resolved by 1% agarose gel electrophoresis, and distinctive bands of approximately 2.3 kb were excised (Figure 1A). The gel-purified dsRNAs were subjected to construct an RNA-Seq library followed by sequencing using Illumina HiSeq 2000. A total of 109 assembled reads with significant fragments per kilobase of transcript per million mapped reads (FPKM) values were obtained. A homology search of the assembled reads suggested the presence of contigs with high similarity to the known viral sequences of RNA-dependent RNA polymerase (RdRp) and capsid protein (CP). Based on the sequence analysis, RT-PCR using the corresponding primer pairs (see Supplementary Table S1) based on two representative contigs (i.e., one for RdRp and the other for CP) resulted in PCR amplicons of the expected sizes, which were subjected to sequence for verification. Northern blot analysis using the dsRNA extracts and probes (see Supplementary Table S1) constructed from PCR amplicons of portions of RdRp and CP revealed a specific hybridization band indicating that the excised dsRNA bands were double bands with similar sizes corresponding each of RdRp and CP. Thus, the analysis of the dsRNA sequences and Northern hybridization indicated that the mycoviral genome is divided into two dsRNA segments (dsRNA1 and dsRNA2) of similar size (Figure 1B). Rapid amplification of cDNA ends (RACE) was conducted to determine the full-length of the dsRNAs.

The complete genome sequences indicated that the dsRNA1 segment was of 2,289 bp with a GC content of 45.1%, and dsRNA2 was of 2,245 bp with a GC content of 49.5%. These sequences were deposited in GenBank (accession number MG973751 and MG973752, respectively). Each segment contains a long open reading frame (ORF) and ORF1 and 2 were on the coding strand of dsRNA1 and dsRNA2, respectively. RACE analysis indicated that the 5′ untranslated region (UTR) of the coding strand of dsRNA1 was 68 nt long and the 3′ UTR was 52 nt long. In addition, the coding strand of dsRNA2 contained a 107-nt-long 5′ UTR and 158-nt-long 3′ UTR. The structure of genome organization of dsRNA1 and dsRNA2 are displayed in Figure 1C.

The sequence of the 5′-termini of both dsRNA1 and 2 (GAACAAGG) were similar to the known consensus sequence of the betapartitivirus GAWWUWNU (N, any nt; W, A or U) and well-matched those of Heterobasidion partitivirus 2 (HetPV2) (Vainio et al., 2011). In addition, characteristic A-rich regions in the 3′-termini were preserved in both dsRNA1 [31 (A) residue in the 3′-terminal 50 nt] and dsRNA2 [48 (A) residue in the 3′-terminal 50 nt]. Like other fungal partitiviruses showing the
characteristics of lengths of 5′- and 3′-non-translated region (NTR), longer 3′-NTR than 5′-NTR was found in dsRNA2 and longer 3′-NTR in dsRNA2 than dsRNA1 was found. Therefore, these results indicated that the obtained sequences were of the full-length of dsRNA1 and 2.

Viral particles were visualized by transmission electron microscopy (TEM). The particles were spherical and of diameter 28–30 nm (Figure 2).

**Phylogenic Analysis of ORF1 and ORF2**

The deduced amino acid sequence of ORF1 consisted of 722 amino acids with a predicted molecular mass of 79.4 kDa. Homology search of the deduced amino acid sequence revealed high similarity to the known sequences of RdRp of Heterobasidion partitivirus 2 (HetPV2) [GenBank accession number, HM565953; E-value, 0; AA identity, 589/722 (81%)]. Phylogenetic analysis using the top ranked similar sequences indicated that our RdRp clustered with the genus Betapartitivirus, and this was supported by significant bootstrap values. We next used the above sequence information for taxonomic reorganization of the family Partitiviridae (Vainio et al., 2015). The extensive analysis showed that our RdRp clustered with HetPV2 and Heterobasidion partitivirus 7 (HetPV7) (Figure 3).

ORF2 was deduced to encode a 659-amino acid protein of predicted molecular mass 72.5 kDa. Homology search showed high similarity to the CP of HetPV2 [GenBank accession number, HM565954; E-value 2e − 65; AA identity, 444/655 (68%)]. Phylogenetic analysis of CP indicated that our sequence clustered strongly to those of HetPV2 and HetPV7. Sequence similarity, genome organization, and phylogenetic analysis indicated that the dsRNAs are genome segments of a new member of the genus Betapartitivirus in the family Partitiviridae. Thus, we named our dsRNA as *Trichoderma harzianum* betapartitivirus 1 (ThPV1).

Based on the ICTV species demarcation criteria for the genus Betapartitivirus in the family Partitiviridae (≤90% and ≤80% amino acid identities in the RdRp and CP, respectively), we conclude that ThPV1 represents a novel species.

**Characteristics of Mycovirus-Cured and -Containing Strains**

Virus-cured isogenic strains were obtained by producing single-spored progenies followed by dsRNA purification. Spores were harvested from 7-day-old culture plates containing actively growing *T. harzianum* NFCF319, and 50 expected colony-forming units (CFUs) were spread on fresh potato dextrose agar (PDA) plate. Of 39 single-spored progenies, ThPV1 dsRNA bands were missing in 31. RT-PCR using gene specific primer pairs did not amplify any band and no hybridization band was observed either, which indicated that ThPV1 was cured in those selected strains. Virus-cured strains were successively sub-cultured weekly and subjected to dsRNA extraction every 4 weeks for 6 months. None of the cured-strains harbored ThPV1 dsRNA, suggesting that they were stable.
To increase the biological replicates, 3 out of 31 cured strains (\textit{T. harzianum} NFCF319-Vf1, -Vf2, and -Vf3), and 3 from 7 vertically transmitted virus-containing clones (\textit{T. harzianum} NFCF319-V1, -V2, and -V3) were randomly selected for further analysis.

No difference in growth rate was observed among the four ThPV1-infected \textit{T. harzianum} isolates, i.e., between the original ThPV1-infected \textit{T. harzianum} NFCF319 strain and its three single-spored virus-containing progenies \textit{T. harzianum} NFCF319-V1, -V2, and -V3. In addition, no significant differences in growth rate or colony morphology were detected between the ThPV1-infected and -cured strains. Although the time of appearance and intensity of the pigmentation varied among individual cultures, a characteristic diffusing yellow pigment developed in all strains during cultivation. In addition, all strains produced similar numbers of gray–green conidia per plate, dispersed in concentric rings (Figure 4).

**Biological Comparison of Mycovirus-Cured and -Containing Strains**

The antagonistic activities of the virus-cured strains against the oyster mushroom, \textit{P. ostreatus}, were measured using culture filtrates of the ThPV1-containing strains (\textit{T. harzianum} NFCF319 and \textit{T. harzianum} NFCF319-V1, -V2, and -V3) and ThPV1-cured strains (\textit{T. harzianum} NFCF319-V1, -V2, and -V3). Significant differences in the growth rate of \textit{P. ostreatus} were observed between medium without and with supplementation with the culture filtrates of \textit{T. harzianum}, regardless of ThPV1 infection. Moreover, significant differences in growth rate were observed between medium supplemented with the culture filtrates of virus-cured and -containing strains. The growth rate was significantly reduced in medium supplemented with culture filtrate of the virus-containing strain (Figure 5 and Figure 6).
Table 1). Colonial growth inhibition was also examined using the plant pathogenic fungus *R. solani* AG-1 (*Figure 6* and *Table 1*). Significant differences in colony growth were observed in supplemented compared with non-supplemented medium. Furthermore, the culture filtrates of the virus-containing strains (*T. harzianum* NFCF319 and *T. harzianum* NFCF319-V1, -V2, and -V3) inhibited the growth of the oyster mushroom and a phytopathogenic fungus more strongly. To examine the effect of the heat-inactivated culture filtrate on fungal growth inhibition, colonial growth of *R. solani* AG-1 was tested using medium supplemented with the heat-inactivated culture filtrate. Colonial growth was inhibited by supplementation with the heat-inactivated culture filtrate, but the effect was not as intense as it was with the original culture filtrate. Interestingly, heat inactivation altered growth inhibition by the culture filtrates of the virus-containing strains to the level of the virus-cured strains.

The activities of two representative antifungal enzymes, β-1,3-glucanase and chitinase, were analyzed. The enzyme activity of the culture filtrate was monitored at 24 h intervals for 4 days. As shown in *Figure 7*, the time course of the β-1,3-glucanase activity was significantly reduced in the culture filtrate of the virus-cured strains of *T. harzianum* NFCF319 (*T. harzianum* NFCF319-Vf denotes the mean of three replicates of each of the NFCF319-Vf, -Vf2, and -Vf3 isolates). Interestingly, there was no difference in chitinase activity between the virus-infected and -cured strains.

To validate the viral effect on the antifungal activity, we performed a “plus” experiment, i.e., we artificially transmitted ThPV1 into a virus-cured strain via anastomosis between virus-infected and -cured strains and analyzed the antifungal activity of the resulting virus-infected recipient strains. First, we transformed the cured strain of *T. harzianum* NFCF319-Vf1 using the hygromycin B resistance cassette to differentiate the ThPV1-transmitted progeny from the parental virus-containing *T. harzianum* NFCF319, because all cured strains were isogenic to their parental strain, i.e., the genetic background of all strains was identical, except for the presence or absence of mycovirus. Among 11 transformants showing stable resistance to hygromycin B after successive transfer to selective medium, the transformant *T. harzianum* NFCF319-Vf1-TNF1 was selected randomly and we observed no phenotypic changes from the parental *T. harzianum* NFCF319-Vf1 strain (*Figure 4*).

After prolonged paired growth of the viral donor (*T. harzianum* NFCF319) and viral recipient (*T. harzianum* NFCF319-Vf1-TNF1) strains, several hyphae at the recipient border were collected and cultured independently on the selective medium, and the presence of ThPV1 was analyzed. Two independent single-spored virus-transmitted hyphal progenies were obtained. One of each single-spored progeny (*T. harzianum* NFCF319-Vf1-TNF1-V1 and -V2) containing ThPV1 was selected for further analysis (*Figure 4*). As shown in *Figures 5, 6*, medium supplemented with the culture filtrates of virus-transmitted *T. harzianum* NFCF319-Vf1-TNF1-V resulted in significant differences in the growth inhibition of *P. ostreatus* and *R. solani*. The level of inhibition was similar to that of the original virus-infected *T. harzianum* NFCF319 and significantly greater than those of virus-cured *T. harzianum* NFCF319-Vf and its transformant *T. harzianum* NFCF319-Vf1-TNF. In addition, significant increases in β-1,3-glucanase activity were observed in *T. harzianum* NFCF319-Vf1-TNF1-V compared with the parental virus-cured transformant *T. harzianum* NFCF319-Vf1-TNF (*Figure 7*). The β-1,3-glucanase activity in *T. harzianum* NFCF319-Vf1-TNF1-V was comparable to that of the original virus-infected *T. harzianum* NFCF319 (*Figure 7*). The enzyme activity profiles of β-1,3-glucanase were similar among the virus-containing strains. The enzyme activity of the virus-containing strains reached a peak at 48 or 72 h and then decreased gradually to 96 h. The enzyme activity profiles of β-1,3-glucanase were similar across the virus-cured strains. However, the levels of the
enzy}m activity of the virus-cured strains were significantly lower at 48 and 72 h of incubation and remained constant until 96 h. No difference in chitinase activity was observed among all strains tested.

DISCUSSION

The two-segmented dsRNA genome (each with one ORF) and isometric particles of 28–30 nm diameter suggest that ThPV1 belongs to the family Partitiviridae. According to the ICTV in 2016, the family Partitiviridae comprises the genera Alphapartitivirus, Betapartitivirus, Gammapartitivirus, Delapartitivirus, and Cryspovirus (Nibert et al., 2014; Lefkowitz et al., 2017). The sizes of the segments and the molecular masses of the encoded proteins of ThPV1 are within the normal range of the genus Betapartitivirus, which are characteristically larger than those of other genera (Nibert et al., 2014). In addition, the high pairwise-identity scores of the amino acid sequences of encoded RdRp (81% to RdRp of HetPV2) and CP (68% to CP of HetPV2) to other known Partitiviridae (Nibert et al., 2014), and the clustering of ThPV1 with other members of the genus Betapartitivirus, indicates that ThPV1 is a new member of the genus Betapartitivirus.

Virus curing by single-spore isolation was successful; i.e., 79.5% of the resulting single-spore colonies were ThPV1-free. In addition, the ThPV1-free proportion was similar (20–84%) among three independent experiments. Therefore, the rate of vertical transmission of ThPV1 to mitotic progenies, conidia, of host fungus was low, although a considerable proportion of conidia showed vertical transmission of ThPV1. Like other mycoviruses, partitiviruses are generally considered to be transmitted vertically during cell division and horizontally during intimate cell-cell contact. Heterobasidion partitiviruses, which are closely related to ThPV1, could be transmitted vertically to basidiospores and conidia and horizontally via hyphal contact (Ihrmark et al., 2002, 2004). The mitotic stability measured by inheritance of mycoviruses during asexual conidiation differed depending on the virus–fungus interaction. The mitotic inheritance of ThPV1 during asexual conidiation was less stable than that of Cryphonectria hypovirus 1 (CHV1) in C. parasitica (Suzuki et al., 2003; Prospero et al., 2006), and of mycoreovirus 1 (MyRV1) in C. parasitica (Sun et al., 2006), but comparable to dsRNAs in the basidiomycetes H. annosum (Ihrmark et al., 2002). The low transmission rate of ThPV1 into conidia is exceptional considering that most vertical transmission of dsRNA into asexual spores of ascomycetes occurs at a markedly higher rate. Indeed, the recently characterized Trichoderma atroviride mycovirus 1 (TaMV1) has a very high transmission rate (33/38) (Lee et al., 2017). Therefore, the low transmission rate of ThPV1 into conidia is attributable to the intrinsic characteristics of the virus–fungus (ThPV1–T. harzianum) interaction.

Considering their persistent lifestyles and direct cell-to-cell transmission, partitiviruses have few, if any, deleterious effects on host cells (Nibert et al., 2014). However, others have reported negative or positive effects of partitiviruses on their host (Suzuki et al., 2001; Márquez et al., 2007; Jenkins et al., 2008; Kanematsu et al., 2010; Vainio et al., 2010, 2017; Hyder et al., 2013; Xiao et al., 2014; Zheng et al., 2014). ThPV1 did not influence the mycelial growth, colony morphology, pigmentation, or conidiation of its host fungus. However, ThPV1 significantly altered the activities of antifungal enzymes potentiating the biocontrol function of its host fungus T. harzianum. It is interesting to see that, regardless of virus-containing or -cured, supplementation of culture filtrates resulted in significantly increased antifungal activities compared to the non-supplementation; furthermore, the heat-treated culture filtrate of the ThPV1-infected strain showed significantly reduced antifungal activities similar to the level of ThPV1-free strain. These results clearly suggested that the antifungal activities were attributable to both heat-stable and -sensitive metabolites. Considering the facts that antifungal

**TABLE 1** Growth inhibition due to the supplementation of culture filtrates.

| Dilution | Strains          | Growth inhibition of R. oryzae (%) | Growth inhibition of R. solani (%) |
|----------|------------------|-----------------------------------|-----------------------------------|
| 1:10     | ThPV1-infected   | 90.29±c                          | 94.78±c                          |
|          | ThPV1-containing | 91.33±c                          | 95.49±c                          |
|          | Isogenic         | 78.82±b                          | 86.87±b                          |
|          | ThPV1-cured      | 81.06±b                          | 86.85±b                          |
|          | ThPV1-free hygromycin B resistant transformant | 91.04±c                          | 95.89±c                          |
| 1:100    | ThPV1-infected   | 82.03±c                          | 87.01±c                          |
|          | ThPV1-containing | 81.64±c                          | 87.56±c                          |
|          | Isogenic         | 66.92±b                          | 80.98±b                          |
|          | ThPV1-cured      | 66.66±b                          | 80.53±b                          |
|          | ThPV1-free hygromycin B resistant transformant | 81.56±c                          | 87.51±c                          |
| 1:1000   | ThPV1-infected   | 66.39±b                          | 78.73±c                          |
|          | ThPV1-containing | 67.06±b                          | 77.08±b                          |
|          | Isogenic         | 66.58±b                          | 5.23±b                           |
|          | ThPV1-cured      | 65.58±b                          | 5.36±b                           |
|          | ThPV1-free hygromycin B resistant transformant | 67.08±b                          | 76.61±c                          |

*ThPV1, Trichoderma harzianum partitivirus 1. Growth inhibition by metabolites from the original ThPV1-infected (T. harzianum NCF319), three single-spored ThPV1-containing (T. harzianum NCF319-V1, -V2, and -V3), isogenic ThPV1-cured (T. harzianum NCF319-VT1, -VT2, and -VT3), ThPV1-free hygromycin B resistant transformant (T. harzianum NCF319-VT1-TNF1-V1 and -V2) isolates. Numbers are the relative inhibition rates (%) compared to without fungal metabolite supplementation. Data were analyzed by ANOVA followed by post hoc Tukey’s with the level of significance at p < 0.05. The same letters at the top of the relative inhibition rate indicate no significant difference between isolates.*
activity was significantly increased by ThPV1, significant increase in the activity of antifungal enzyme of β-1,3-glucanase was observed in the ThPV1-infected strain, and heat treatment reversed the increased antifungal activity of ThPV1-infected strain, the antifungal activity was likely mediated by a hydrolytic enzyme, such as β-1,3-glucanase. ThPV1 modulates the activity of β-1,3-glucanase but not that of chitinase. These results suggest that ThPV1 influences the activity of a specific fungal enzyme, which enhances the mycoparasitism of ThPV1-infected T. harzianum. Mycoviruses can exert complex effects on their fungal host, which mediated osmotic stress tolerance (Nerva et al., 2017) and mycotoxin accumulation (Nerva et al., 2018). Thus, whether ThPV1 exerts antagonistic effects on other fungal strains or species should be the subject of further studies.

In addition to the curing experiment including multiple virus-cured and -retained progenies, horizontal viral transmission via hyphal fusion allowed us to verify that the changes in antifungal activities were due to ThPV1. The fact that, compared with the original T. harzianum NFCF319 strain, all the viral-transmitted isolates showed similar levels of antifungal activity, such as the growth inhibition of other fungi, indicates that ThPV1 is responsible for the enhanced antifungal activity of the host fungus. Although further work is required, it is important to note the increased β-1,3-glucanase activity is implicated in the hydrolysis of pathogenic fungi during mycoparasitic attack by T. harzianum (Qualhato et al., 2013). The antifungal activity was higher in the culture filtrate of ThPV1-infected strains, suggesting that the enhanced β-1,3-glucanase activity influences the potential antifungal activity of ThPV1-infected strains.

To our knowledge, this is the first report of a betapartitivirus in T. harzianum. In addition, the mycovirus enhanced the antifungal activity of the host fungus by regulating the activity of a specific antifungal enzyme.

**AUTHOR CONTRIBUTIONS**

D-HK supervised the experiments. JC and H-EY performed the experiments. D-HK, JC, and H-EY prepared the figures and edited the manuscript. D-HK wrote the manuscript.

**FUNDING**

This work was supported by the Bio-industry Technology Development Program, Ministry for Food, Agriculture, Forestry and Fisheries, Republic of Korea and in part by the NRF grant by MSIP (2018R1A2A1A05078682). We thank the Institute of Molecular Biology and Genetics at Chonbuk National University for kindly providing the facilities for this research. D-HK was supported by “Research Base Construction Fund Support Program” funded by Chonbuk National University in 2016.

**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2018.01699/full#supplementary-material

**REFERENCES**

Churchill, A. C. L., Ciuffetti, L. M., Hansen, D. R., Van Ettten, H. D., and Van Alfen, N. K. (1990). Transformation of the fungal pathogen Cryptocnemtria parasitica with a variety of heterologous plasmids. Curr. Genet. 17, 25–31. doi: 10.1007/BF03313245

Cullen, D., Leong, S. A., Wilson, L. J., and Henner, D. J. (1987). Transformation of Aspergillus nidulans with the hygromycin-resistance gene, hph. Gene 57, 21–26. doi: 10.1016/0378-1119(87)90172-7

Fitch, W. M. (1971). Toward defining the course of evolution: minimum change for a specific tree topology. Syst. Zool. 20, 406–416. doi: 10.2307/2412116

Ghabrial, S. A. (1998). Origin, adaptation and evolutionary pathways of fungal viruses. Virus Genes 16, 119–131. doi: 10.1023/A:1007966229995

Ghabrial, S. A., Caston, J. R., Jiang, D., Nibert, M. L., and Suzuki, N. (2015). 50-plus years of fungal viruses. Virology 47, 356–368. doi: 10.1016/j.virology.2015.02.034

Hatvani, N., Kredics, L., Antal, Z., and Mész, I. (2002). Changes in activity of extracellular enzymes in dual cultures of Lentinula edodes and mycoparasitic trichoderma strains. J. Appl. Microbiol. 92, 415–423. doi: 10.1046/j.1365-2672.2002.01542.x

Howell, C. R. (2003). Mechanisms employed by trichoderma species in the biological control of plant diseases: the history and evolution of current concepts. Plant Dis. 87, 4–10. doi: 10.1094/PDIS.2003.87.1.4

Hyder, R., Pennanen, T., Hamberg, L., Vainio, E. J., Piri, T., and Hantula, J. (2013). Two viruses of Heterobasidion confer beneficial, cryptic or detrimental effects to their hosts in different situations. Fungal Ecol. 6, 387–396. doi: 10.1016/j.funeco.2013.05.005

Ihrmark, K., Johannesson, H., Stenström, E., and Stenlid, J. (2002). Transmission of double-stranded RNA in Heterobasidion annosum. Fungal Genet. Biol. 36, 147–154. doi: 10.1016/S1087-1845(02)00011-7

Ihrmark, K., Stenström, E., and Stenlid, J. (2004). Double-stranded RNA transmission through basidiospores of Heterobasidion annosum. Mycol. Res. 108, 149–153. doi: 10.1017/S0953756203008839

Jenkins, M. C., Higgins, J., Abrahante, J. E., Kniel, K. E., O’Brien, C., Trout, J., et al. (2008). Fecundity of Cryptosporidium parvum is correlated with intracellular levels of the viral symbiont CPV. Int. J. Parasitol. 38, 1051–1055. doi: 10.1016/j.ijpara.2007.11.005

Kanematsu, S., Sasaki, A., Onoue, M., Oikawa, Y., and Ito, T. (2010). Extending the fungal host range of a partitivirus and a mycoreovirus from Rosellinia necatrix by inoculation of protoplasts with virus particles. Phytopathology 100, 922–930. doi: 10.1094/PHYTO-100-9-0922

Kim, J. M., Jung, J. E., Park, J. A., Park, S. M., Cha, B. J., and Kim, D. H. (2013). Biological function of a novel chrysovirus, CnV1-BS122, in the korean cryphonectria nitchiki BS122 strain. J. Biocoi. Bioeng. 115, 1–3. doi: 10.1016/j.jbiosc.2012.08.007

Kim, M. J., Choi, J. W., Park, S. M., Cha, B. J., Yang, M. S., and Kim, D. H. (2002). Characterization of a fungal protein kinase from Cryptocnemtria parasitica and its transcriptional upregulation by hypovirus. Mol. Microbiol. 45, 933–941. doi: 10.1046/j.1365-2958.2002.03079.x

Kredics, L., García Jimenez, L., Naeimi, S., Czifra, D., Urbán, P., Manczinger, L., et al. (2010). “A challenge to mushroom growers: the green mould disease of cultivated champignons,” in Current Research, Technology and Education Topics in Applied Microbiology and Microbial Biotechnology, ed. A. Méndez-Vilas (Badajoz: Formatex), 295–305.

Kucizek, C. P., and Penttila, M. E. (1998). “Regulation of production of plant polysaccharide degrading enzymes by Trichoderma,” in Trichoderma and Gliocladium: Enzymes, Biological Control and Commercial Applications, eds G. E. Harman and C. P. Kucizek (London, UK: Taylor and Francis), 49–71.
Kumar, S., Stecher, G., and Tamura, K. (2016). MEGA7: molecular evolutionary genetics analysis version 7.0 for bigger datasets. Mol. Biol. Evol. 33, 1870–1874. doi: 10.1093/molbev-msw054

Lee, S. H., Yun, S. H., Chun, J., and Kim, D. H. (2017). Characterization of a novel dsRNA mycovirus of Trichoderma atroviride NFC028. Arch. Virol. 162, 1071–1077. doi: 10.1007/s00705-016-2314-2

Leikowitz, E. J., Adams, M. J., Davison, A. J., Siddell, S. G., and Simmonds, P. (2017). Virus Taxonomy: Classification and Nomenclature of Viruses. The Online 10th Report of the ICTV. Available at: https://talk.ictvonline.org/ictv-reports/ictv_online_report

Márquez, L. M., Redman, R. S., and Roossinck, M. J. (2007). A virus in a fungus in a plant: three-way symbiosis required for thermal tolerance. Science 315, 513–515. doi: 10.1126/science.1136237

Nerva, L., Chitarra, W., Siciliano, I., Gaiotti, F., Ciuffo, M., Forgia, M., et al. (2018). Significance of dsDNA genetic elements in plant pathogenic fungi. Ann. Rev. Phytopathol. 38, 37–55. doi: 10.1146/annurev.phyto.38.050213.094315

Nibert, M. L., Ghabrial, S. A., Maiss, E., Lesker, T., Vainio, E. J., Jiang, D., et al. (2014). Taxonomic reorganization of family Partitiviridae and other recent progress in partitivirus research. Virus Res. 188, 128–141. doi: 10.1016/j.virusres.2014.04.007

Nuss, D. L. (2005). Hypovirulence: mycoviruses at the fungal-plant interface. Nat. Rev. Microbiol. 3, 632–642. doi: 10.1038/nrmicro1206

Nuss, D. L., and Koltin, Y. (2019). Significance of dsDNA genetic elements in plant pathogenic fungi. Annu. Rev. Phytopathol. 28, 37–55. doi: 10.1146/annurev.phyto.28.090190.000345

Papavizas, G. C. (1985). Trichoderma and Gliocladium: biology, ecology, and potential for biocontrol. Ann. Rev. Phytopathol. 23, 23–54. doi: 10.1146/annurev.phyto.23.091805.000323

Park, S. M., Kim, J. M., Chung, H. J., Lim, J. Y., Kwon, B. R., Lim, J. G., et al. (2008). Occurrence of diverse dsRNA in a korean population of the chestnut blight fungus, Cryphonectria parasitica. Mycol. Res. 112, 1220–1226. doi: 10.1016/j.mycres.2008.04.001

Penttilä, M., Nevalainen, H., Rättö, M., Salminen, E., and Knowles, J. (1987). A versatile transformation system for the cellulolytic filamentous fungus Trichoderma reesei. Gene 51, 155–164. doi: 10.1016/0378-1119(87)90110-7

Prospero, S., Conedera, M., Heinerig, U., and Rigling, D. (2006). Saprophytic activity and sporulation of Cryphonectria parasitica on dead chestnut wood in forests with naturally established hypovirus. Phytopathology 96, 1337–1344. doi: 10.1094/PHYTO-96-1337

Qualhato, F. T., Lopes, F. A. C., Steinidorff, A. S., Brandão, R. S., Jesuino, R. S. A., and Ulhoa, C. J. (2013). Mycoparasitism studies of trichoderma species against Rhizoctonia solani, “in Trichoderma and Gliocladium: Basic Biology, Ecology, and Agricultural Applications,” ed. G. E. Harman and C. P. Kubicek (London, UK: Taylor and Francis), 267–288.

Sivasithamparam, K., and Ghisalberti, E. L. (1998). “Secondary metabolism in Trichoderma and Gliocladium,” in Trichoderma and Gliocladium: Basic Biology, Taxonomy and Genetics, eds G. E. Harman and C. P. Kubicek (London, UK: Taylor and Francis), 139–191.

Sun, L., Nuss, D. L., and Suzuki, N. (2006). Mycoviruses mediate mycoytoxin regulation in Aspergillus ochraceus. Environ. Microbiol. 315, 513–515. doi: 10.1126/science.1136237

Van Alfen, N. K. (1986). “Hypovirulence of Endothia (Cryphonectria) parasitica and Rhizoctonia solani,” in Fungal Virology, ed. K. W. Buck (Boca Raton, FL: CRC Press), 143–162.

Vainio, E. J., Korhonen, K., Tuomivirta, T. T., and Hantula, J. (2010). A novel putative partitivirus of the saprotrophic fungus Heterobasidion annosum P-type partitivirus. Arch. Virol. 155, 79–86. doi: 10.1007/s00705-010-0823-9

Vainio, E. J., Jurvansuu, J., Hyder, R., Kashif, M., Piri, T., and Tuomivirta, T. (2017). The partitivirus HетPV13-an1 mediates severe growth debilitation and major alterations in the gene expression of a fungal forest pathogen. J. Virol. 92,e01744-17. doi: 10.1128/JVI.01744-17

Vainio, E. J., Kerio, S., and Hantula, T. (2011). Description of a new putative virus infecting the conifer pathogenic fungus Heterobasidion annosum P-type partitivirus. Arch. Virol. 156, 985–988. doi: 10.1007/s00705-010-0823-9

Wickner, R. B. (1992). Double-stranded and single-stranded RNA viruses of Rhizobium leguminosarum by the nod genes derived from Rhizobium leguminosarum bv. trifolii strain 4S. Gene 245, 77–84. doi:10.1016/0378-1119(89)80377-8

Xiao, X., Cheng, J., Tang, J., Fu, Y., Jiang, D., Baker, T. S., et al. (2014). A novel partitivirus that confers hypovirulence on plant pathogenic fungi. J. Virol. 88, 10120–10133. doi: 10.1128/JVI.01036-14

Yun, S. H., Lee, S. H., So, K. K., Kim, J. M., and Kim, D. H. (2016). Incidence of diverse dsRNA mycoviruses in trichoderma spp. causing green mold disease of shiitake Lentinula edodes. FEMS Microbiol. Lett. doi:10.1093/femsle/fnvw220 [Epub ahead of print].

Zheng, L., Zhang, M., Chen, Q., Zhu, M., and Zhou, E. (2014). A novel mycovirus closely related to viruses in the genus Alphapartitivirus confers hypovirulence in the phytopathogenic fungus Rhizoctonia solani. Virol. 456-457, 220–226. doi: 10.1016/j.virology.2014.05.029

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

Copyright © 2018 Chun, Yang and Kim. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.