Metabolomics Investigation of an Association of Induced Features and Corresponding Fungus during the Co-culture of *Trametes versicolor* and *Ganoderma applanatum*

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The co-culture of *Trametes versicolor* and *Ganoderma applanatum* is a model of intense basidiomycete interaction, which induces many newly synthesized or highly produced features. Currently, one of the major challenges is an identification of the origin of induced features during the co-culture. Herein, we report a 13C-dynamic labeling analysis used to determine an association of induced features and corresponding fungus even if the identities of metabolites were not available or almost nothing was known of biochemical aspects. After the co-culture of *T. versicolor* and *G. applanatum* for 10 days, the mycelium pellets of *T. versicolor* and *G. applanatum* were sterilely harvested and then mono-cultured in the liquid medium containing half fresh medium with 13C-labeled glucose as carbon source and half co-cultured supernatants collected on day 10. 13C-labeled metabolome analyzed by LC-MS revealed that 31 induced features including 3-phenyllactic acid and orsellinic acid were isotopically labeled in the mono-culture after the co-culture stimulation. Twenty features were derived from *T. versicolor*, 6 from *G. applanatum*, and 5 features were synthesized by both *T. versicolor* and *G. applanatum*. 13C-labeling further suggested that 12 features such as previously identified novel xyloside [N-(4-methoxyphenyl)formamide 2-O-beta-D-xyloside] were likely induced through the direct physical interaction of mycelia. Use of molecular network analysis combined with 13C-labeling provided an insight into the link between the generation of structural analogs and producing fungus. Compound 1 with m/z 309.0757, increased 15.4-fold in the co-culture and observed 13C incorporation in the mono-culture of both *T. versicolor* and *G. applanatum*, was purified and identified as a phenyl polyketide, 2,5,6-trihydroxy-4,
6-diphenylcyclohex-4-ene-1,3-dione. The biological activity study indicated that this compound has a potential to inhibit cell viability of leukemic cell line U937. The current work sets an important basis for further investigations including novel metabolites discovery and biosynthetic capacity improvement.

**Keywords:** basidiomycete, interaction, $^{13}$C-labeling, origin of induced features, phenyl polyketide, metabolite identification, LC-MS

## INTRODUCTION

Secondary metabolites are an important source of valuable drug leads, of which compounds derived from various fungi, especially medicinal fungi of basidiomycetes, represent an important part. To explore chemical diversity, several approaches such as epigenetic modification or non-targeted metabolic pathway manipulation have recently been developed in the *Aspergillus* and *Streptomyces* species (Scherlach and Hertweck, 2009; Chiang et al., 2011). In the case of basidiomycetes, the induction of novel secondary metabolites or enhancement of secreting extracellular enzymes can be achieved by activating cryptic biosynthetic pathways through establishment of a fungi interaction in the co-culture (Peiris et al., 2008; Hiscox et al., 2010). This co-culture strategy mimics natural ecosystem, in which interspecies interaction of basidiomycetes is very common (Hiscox et al., 2015, 2017). Recently, Zheng et al. demonstrated that co-culture of basidiomycetes *Inonotus obliquus* and *Phellinus punctatus* resulted in the accumulation of lanostane-type triterpenoids, polyphenols, and melanins, compounds capable of scavenging free radicals and inhibiting tumor cell proliferation (Zheng et al., 2011). The co-culture of *Trichoderma Reesi* with *Coprinus comatus* was an important approach for an on-site generation of lignocellulolytic enzyme leading to the increase of lignocellulose degradation rate as described by Ma and Ruan (2015). In our previous work, fifteen wood-decaying basidiomycetes and two straw-decaying basidiomycetes were used to establish 136 pairwise co-cultures on agar plates (Yao et al., 2016). The co-culture system of *Trametes versicolor* and *Ganoderma applanatum* showed an interaction zone, in which the accumulation of a series of known carboxylic acids as well as novel xylosides were observed.

The application of co-culture has enhanced the number of discovered novel secondary metabolites but also raises several challenges. One of the major challenges is how to timely and accurately identify the species responsible for the newly induced metabolites during the microorganism interaction. Typically, the structure of compounds and related biochemical information were required to verify the unique secondary metabolite and its particular producer (Riedlinger et al., 2006). Recently, a combined technique of nanospray desorption electrospray ionization and matrix-assisted laser desorption/ionization-time of flight mass spectrometry has been developed as a platform to reveal many unknown metabolites produced by *Streptomyces coelicolor* when paired with five other actinomycetes (Traxler et al., 2013). However, this approach was limited to associate the novel metabolites with corresponding producer grown on agar plates and was not suitable to the co-culture in the liquid medium. The metabolic incorporation of stable isotopes $^{13}$C or $^{15}$N is a powerful approach used for quantitative proteome and metabolome studies (Julka and Regnier, 2004; Yang et al., 2010). Moreover, $^{13}$C-dynamic labeling analysis has been applied for analyzing the metabolite turnover rates, distinguishing the flux distribution between two pathways starting from the same metabolic point, and interpreting the synthetic process of novel metabolites (Yang et al., 2012; Shlomi et al., 2014; Hammerl et al., 2017). With the advantage of a $^{13}$C-labeling approach, the current study reports on the identification of the producer of novel induced metabolites in liquid medium after the two interactive mycelia were sterilely separated and mono-cultured after the induction by the co-culture.

The research of the interspecies crosstalk expands our possibilities to discover novel metabolites, and also increases our understanding of how these metabolites are induced in microbial consortia. A chemical warfare in the fungal-fungal communication is often described as diffusion of harmful and chemically complex metabolites from one partner to the other (Bertrand et al., 2014). For instance, the interaction between the *Aspergillus niger* and *Aspergillus flavus* led to the inhibition of aflatoxin B1 produced by *A. flavus* through signal molecules downregulating expression of major biosynthetic genes (Xing et al., 2017). Under some conditions, co-cultivation appears to trigger production and accumulation of novel metabolites without the involvement of released signaling molecules. For instance, co-cultivation with *Corynebacteria glutamicum* or *Tsukamurella pulmonis* was indicated to stimulate a novel pathway in *S. endus*, contributing to a new heterocyclic chromophore-containing antibiotics alchivemycin A (Onaka et al., 2011). Notably, the mono-culture of *S. endus* did not generate the same compound with or without the addition of filter sterilized supernatants from bacterial culture. The production of alchivemycin A therefore appeared to require a direct physical interaction between *S. endus* and the coryneform bacteria cells. For this research, $^{13}$C dynamic labeling was further utilized to suggest the potential mechanism of the induction of increased and newly synthesized features.

In this work, we built upon the characteristic of mycelial pellets of basidiomycetes and $^{13}$C-labeling analysis, to analyze and catalog a broad range of $^{13}$C-labeled features, which were highly accumulated in the co-culture of *T. versicolor* and *G. applanatum*. $^{13}$C dynamic labeling further suggested two potential mechanism of the induction of these features. Ultimately, compound 1 with M+H$^+$ m/z 309.0757, that displayed $^{13}$C incorporation in the mono-culture of both *T. versicolor* and *G. applanatum*, was isolated and identified as novel phenyl polyketide, and its biological activity was evaluated.
MATERIALS AND METHODS

Chemicals
All chemicals including standards (ascorbic acid and phenolic antioxidant 2, 6-di-tert-butyl-4-methoxyphenol) were purchased from either Sigma-Aldrich (St. Louis, MO, USA) or TCI (Kita-ku, Tokyo, Japan). Millipore water (Billerica, MA, USA) was used for the preparation of all the media and sample solutions.

Fungi Material and Culture Conditions
G. applanatum (CGMCC No. 5,249) and T. versicolor (CGMCC No. 12241) were deposited at the Shandong Province Key Lab of Applied Mycology in China. The culture medium was supplied at concentrations as follows: 2 g glucose, 0.2 g KH₂PO₄, 0.1 g MgSO₄, 0.4 g peptone, and 4 g agar (only in solid medium) in 200 mL of sterilized water.

Mono-culture of T. versicolor and G. applanatum on Agar Plate
The mono-culture procedure was adapted from our previous publication (Yao et al., 2016), and mono-culture medium was the same as above. Briefly, a 5 mm agar plug of each fungus scraped from agar slant culture-medium was cultured on a Petri dish (9 cm diameter), and were incubated at 28°C for 10 days.

Co-culture of T. versicolor and G. applanatum in Liquid Medium and Sample Preparation
Eight 5 mm agar plugs of T. versicolor and G. applanatum were separately pre-cultured in 500 mL shake flask containing 200 mL of culture medium at 28°C for 4 days on orbital shakers at 180 rpm. Then 100 mL culture broth of G. applanatum was transferred into the culture of 100 mL T. versicolor, and co-cultivated up to 18 days. All the co-cultures had three independent biological replicates. At the harvest time, 10 mL of co-culture broth was filtered by using MILLEX-GP PES membrane filters (0.22 µm, 33 mm, Merck Millipore, Germany) and the filtrate was dried in a freeze-dryer ALPHA 1–2LD plus (Christ, Osterode, Germany). Five milliliters of freshly prepared dichloromethane/methanol/water (64:36:8, v/v) solvent mixture was added to the dried samples (Yao et al., 2016). The sample extractions were carried out in a water bath sonicator (KQ-300GVDV, Kunshan, China) at 25°C for 20 min, and were centrifuged at 12,000 rpm for 10 min. Finally, the extracts were dried on a rotational vacuum concentrator (Christ, Osterode, Germany) and stored in a −80°C freezer.

Measurement of the Metabolome
The extracts were dissolved in 200 µL methanol, and then were centrifuged at 12,000 rpm for 15 min. The supernatants were transferred into 250 µL Agilent autosampler vials. The samples were analyzed on an Agilent liquid chromatograph-quadrupole time-of-flight mass spectrometer (LC-QTOF-MS, Agilent 1290 Infinity-6530B, Agilent Technologies, Santa Clara, CA, USA) as previously described (Cui et al., 2016; Hu et al., 2016). Briefly, 10 µL of the samples was separated on an Acquity UPLC BEH C18 column (100 × 2.1 mm, 1.7 µm, Waters, Milford, MA, USA). The mobile phase A was water with 0.1% formic acid. The mobile phase B was pure acetonitrile. The reversed-phase liquid chromatographic elution gradient was optimized in order to maximize the resolution of the induced features. The gradient was the following: 0–3 min, 5% B, 3–12 min, 5–40% B, 12–38 min, 40–95% B, 38–46 min, 95% B, 46–48 min, 95–5% B, 48–55 min, 5% B. This shallow gradient provided reproducible separation for many co-eluting compounds in a time window from 5.0 to 8.0 min. The TOF m/z range was set to 50–1,200 amu in centroid mode with a scan rate of 1.5 spectra/s. All the samples had three independent biological replicates. Each biological replicate had two analytical replicates.

Data Pre-processing and Principal Component Analysis
LC-QTOF-MS data were converted into mzML format using MS Convert software (Holman et al., 2014). Data pre-processing and statistical analysis were performed with MZmine 2 (Version 2.11) (Pluskal et al., 2010) and SIMCA-P 11.5. For the MZmine 2, the peak detection threshold for MS signal intensity was set to 1.0 × 10³. The chromatogram building was realized using a minimum time span of 0.01 min, minimum height of 2.5 × 10³, and m/z tolerance of 0.005 (or 10 ppm). Chromatograms were deconvoluted with the following settings: search minimum in absolute retention time (RT) range 0.1 min, minimum relative height 10%, minimum absolute height 2.5 × 10³ and baseline level 1.2. The chromatogram isotopic peaks grouper algorithm was set as m/z tolerance of 0.005 (or 10 ppm) and absolute RT tolerance of 0.10 min. Chromatograms were peak aligned with m/z tolerance at 0.008 (or 15 ppm) and absolute RT tolerance 1 min. The peak list was eventually gap-filled with m/z tolerance at 0.008 (or 15 ppm), and absolute RT tolerance of 0.20 min. To classify m/z in the peak list, principal component analysis (PCA) was carried out by using SIMCA-P (version 11.5). Two steps were required to perform PCA. The first step was to set the variables m/z and RT as Primary ID and Secondary ID. The secondary step was to do the normalization of the variables with Pareto scaling. This normalized method was embedded in SIMPCA-P, which did not require further parameter tuning. After that, PCA were displayed by a scores plot, mainly observing the overall cluster of the different treatments as well as the presence of outliers. The correlation coefficient loading plot was used to identify the variables responsible for the clustering or separation of the treatments.

Molecular Network Analysis
MS/MS data for molecular network analysis were acquired in targeted MS/MS mode on the same LC-QTOF-MS system (the precursor ions are listed in Table 1). The collision energy and m/z range for different precursor ions were optimized based on their own characteristics as our previous publication (Yao et al., 2016). MS/MS data were converted to mzML format, and then were subjected to the Molecular Networking workflow of Global Natural Products Social Molecular (GNPS at gnps.ucsd.edu) using the Group Mapping feature (Wauters et al., 2012; Wang et al., 2016). The subnetworks were generated with settings of minimum pairs cosine 0.65, parent mass tolerance 1.0 Dalton,
| m/z(–) | RT /min | Predictive elemental composition | Difference | Produced by fungus | 12C signal intensity in the co-culture | Features discovered with the current, optimized chromatography |
|--------|---------|---------------------------------|------------|-------------------|--------------------------------------|--------------------------------------------------|
| 136.0403 | 3.91 | C7H7NO2 | Newly synthesized | G. applanatum | 4.00E+04 | |
| 165.0554 | 9.79 | C9H17O3 | Fold increase 15.8 ± 0.7 | T. versicolor and G. applanatum | 2.50E+04 | |
| 166.0507 | 8.65 | C9H9NO3 | Newly synthesized | NA | 6.00E+04 | |
| 167.0348 | 9.90 | C9H9O4 | Fold increase 5.8 ± 0.6 | G. applanatum | 5.00E+03 | |
| 181.0815 | 6.46 | C9H11N2O3 | Newly synthesized | T. versicolor | 4.00E+03 | +
| 195.0505 | 8.37 | C9H12O4 | Fold increase 25.3 ± 1.2 | NA | ND | |
| 244.0612 | 6.86 | C13H11NO4 | Newly synthesized | NA | 7.00E+03 | |
| 249.1497 | 16.33 | C15H22O3 | Newly synthesized | T. versicolor | 1.50E+04 | +
| 251.0712 | 15.72 | C16H23O3 | Fold increase 7.3 ± 1.3 | T. versicolor | 6.00E+03 | +
| 269.0577 | 11.75 | C14H10N2O4 | Newly synthesized | G. applanatum | 9.00E+03 | +
| 271.0716 | 7.03 | C14H12N2O4 | Newly synthesized | G. applanatum | 1.00E+05 | |
| 271.0725 | 10.25 | C14H12N2O4 | Newly synthesized | G. applanatum | 7.00E+04 | |
| 279.0666 | 13.25 | C17H12O4 | Fold increase 5.2 ± 1.3 | T. versicolor | 2.50E+05 | +
| 281.0808 | 11.93 | C17H14O4 | Fold increase 102.2 ± 4.8 | T. versicolor | 5.00E+04 | |
| 296.0928 | 8.65 | C13H17NO7 | Newly synthesized | NA | 2.30E+04 | |
| 304.0626 | 3.33 | C9H13N2O9 | Newly synthesized | NA | ND | |
| 306.0775 | 10.46 | C18H13NO4 | Newly synthesized | NA | ND | + |
| 309.0756 | 12.04 | C18H14O5 | Fold increase 15.4 ± 1.2 | T. versicolor and G. applanatum | 2.50E+05 | +
| 311.0931 | 11.49 | C18H16O5 | Fold increase 400 ± 7.1 | T. versicolor | 4.00E+04 | |
| 314.0899 | 3.32 | C12H13N2O4 | Newly synthesized | NA | ND | |
| 330.2666 | 14.03 | C19H17NO4 | Fold increase 59.7 ± 8.1 | T. versicolor | 6.00E+04 | |
| 334.0733 | 11.44 | C20H17N2O4 | Newly synthesized | NA | ND | |
| 337.0712 | 10.59 | C20H19N2O2 | Newly synthesized | NA | ND | + |
| 341.1046 | 16.16 | C20H14N2O2 | Newly synthesized | T. versicolor | 5.00E+03 | +
| 342.2664 | 15.17 | C19H17NO4 | Newly synthesized | T. versicolor and G. applanatum | 6.00E+04 | |
| 387.1926 | 20.12 | C19H18N2S3 | Newly synthesized | NA | ND | |
| 389.2072 | 20.07 | C19H18N2S3 | Newly synthesized | NA | ND | |
| 397.223 | 19.42 | C21H34O7 | Newly synthesized | NA | 1.25E+04 | |
| 398.1058 | 13.53 | C20H16N2O4 | Newly synthesized | NA | ND | + |
| 399.2396 | 20.07 | C21H32O7 | Fold increase 103.3 ± 4.2 | NA | 3.00E+03 | |
| 405.0737 | 12.83 | C22H10N2O3 | Newly synthesized | NA | ND | |
| 406.1043 | 7.89 | C21H13N2O2 | Newly synthesized | NA | ND | |
| 415.2338 | 10.40 | C21H18O6 | Newly synthesized | NA | ND | |
| 436.2698 | 15.86 | C22H16N6O6 | Newly synthesized | T. versicolor | 2.50E+04 | +
| 454.2817 | 15.86 | C24H15NO7 | Newly synthesized | T. versicolor | 1.50E+04 | +
| 455.1685 | 8.64 | C21H12N6O6 | Newly synthesized | NA | ND | |
| 472.2293 | 15.86 | C24H15NO8 | Newly synthesized | T. versicolor | 1.25E+04 | +
| 490.3025 | 15.06 | C24H15NO9 | Newly synthesized | T. versicolor | 3.40E+04 | +
| 567.3139 | 15.87 | C25H18N2O12 | Newly synthesized | T. versicolor | 7.00E+03 | +
| 581.1206 | 13.24 | C26H22O8 | Newly synthesized | NA | ND | +
| 586.2857 | 16.44 | C28H14N12O6 | Newly synthesized | NA | 2.00E+03 | +
| 597.1968 | 8.63 | C31H34O12 | Newly synthesized | NA | ND | |
| 629.1419 | 11.77 | C30H22N4O12 | Newly synthesized | NA | ND | |
| 641.1461 | 12.65 | C19H19N2O18 | Newly synthesized | NA | ND | |
| 138.0555 | 3.93 | C7H7NO2 | Newly synthesized | G. applanatum | 8.00E+04 | |

(Continued)
ion tolerance 0.5 Dalton, maximum connected components 50, minimum matched peaks 6, minimum cluster size 2. The results were then visualized using Cytoscape (Version 3.1.1) (Su et al., 2014).

**13C-Labeling Analysis**

T. versicolor and G. applanatum were co-cultivated up to 10 days as described above. Then mycelium pellets of T. versicolor and G. applanatum were respectively harvested using sterilized tweezers based on the diameter size and color. The mycelium pellets of T. versicolor had the size ranging from 6 to 8 mm with the color of faint yellow and those of G. applanatum had the size ranging from 2 to 4 mm with red color. The harvested mycelium pellets were washed with sterile water three times and then mono-cultured in 50 mL shake flask containing 10 mL fresh medium with $^{13}$C-labeled glucose at the final concentration of $5 \text{ g L}^{-1}$ and 10 mL of co-cultured supernatants collected on day 10. For the control experiment no co-culture supernatants

| m/z(+) | RT / min | Predictive elemental composition | Difference | Produced by fungus | $^{13}$C signal intensity in the co-culture | Features discovered with the current, optimized chromatography |
|--------|----------|----------------------------------|------------|-------------------|------------------------------------------|----------------------------------------------------------|
| 140.0708 | 8.64     | C$_7$H$_7$NO$_2$ | Fold increase $14.9 \pm 1.3$ | G. applanatum | 1.0E+04 | |
| 150.0548 | 8.53     | C$_8$H$_7$NO$_2$ | Newly synthesized | NA | 1.0E+05 | |
| 160.0377 | 3.84     | C$_6$H$_5$N$_2$O$_4$ | Newly synthesized | NA | ND | |
| 168.0653 | 8.65     | C$_7$H$_8$NO$_3$ | Newly synthesized | NA | 8.0E+04 | +
| 196.0944 | 9.16     | C$_{10}$H$_{13}$NO$_3$ | Newly synthesized | NA | 2.0E+03 | +
| 216.1021 | 14.2     | C$_{13}$H$_{14}$NO$_2$ | Newly synthesized | NA | 1.0E+04 | +
| 230.1177 | 19.07    | C$_{14}$H$_{15}$NO$_2$ | Newly synthesized | NA | 2.6E+03 | |
| 257.0919 | 9.18     | C$_{14}$H$_{15}$NO$_3$ | Newly synthesized | NA | ND | +
| 265.0860 | 11.92    | C$_{17}$H$_{22}$O$_3$ | Fold increase $802.4 \pm 9.6$ | NA | ND | +
| 268.2277 | 15.96    | C$_{16}$H$_{19}$N$_2$O$_2$ | Newly synthesized | NA | 4.0E+03 | +
| 270.0979 | 3.27     | C$_{13}$H$_{11}$N$_2$O$_2$ | Newly synthesized | NA | ND | |
| 272.1129 | 3.58     | C$_{12}$H$_{11}$N$_2$O$_2$ | Newly synthesized | NA | ND | |
| 273.0871 | 10.22    | C$_{14}$H$_{12}$N$_2$O$_4$ | Newly synthesized | G. applanatum | 1.1E+05 | |
| 286.1436 | 14.85    | C$_{17}$H$_{19}$NO$_3$ | Newly synthesized | NA | ND | |
| 287.1030 | 10.61    | C$_{16}$H$_{11}$N$_2$O$_4$ | Newly synthesized | NA | ND | |
| 289.2900 | 20.88    | C$_{17}$H$_{23}$N$_2$O$_2$ | Fold increase $50.5 \pm 3.3$ | T. versicolor | 1.9E+04 | +
| 298.2744 | 18.56    | C$_{18}$H$_{25}$NO$_2$ | Newly synthesized | T. versicolor | 4.0E+04 | +
| 300.1076 | 8.65     | C$_{13}$H$_{17}$NO$_7$ | Newly synthesized | NA | 3.0E+04 | |
| 302.3069 | 21.38    | C$_{18}$H$_{19}$N$_2$O$_3$ | Fold increase $24.4 \pm 3.1$ | T. versicolor and G. applanatum | 4.8E+05 | |
| 322.0898 | 8.7      | C$_{11}$H$_{11}$N$_2$O$_5$ | Newly synthesized | NA | ND | +
| 330.3371 | 26.77    | C$_{20}$H$_{12}$N$_2$O$_2$ | Fold increase $152.3 \pm 1.6$ | T. versicolor | 5.6E+03 | |
| 332.8280 | 14.05    | C$_{18}$H$_{23}$NO$_3$ | Fold increase $74.3 \pm 4.2$ | T. versicolor | 3.6E+04 | +
| 334.9365 | 14.95    | C$_{18}$H$_{59}$NO$_4$ | Newly synthesized | T. versicolor | 1.6E+04 | +
| 344.2503 | 15.17    | C$_{19}$H$_{31}$NO$_4$ | Newly synthesized | T. versicolor | 1.25E+05 | +
| 348.1430 | 11.71    | C$_{20}$H$_{27}$N$_2$O$_3$ | Newly synthesized | NA | ND | |
| 358.2965 | 15.66    | C$_{20}$H$_{59}$NO$_4$ | Fold increase $25.2 \pm 2.2$ | T. versicolor | 3.0E+04 | +
| 369.2269 | 13.47    | C$_{16}$H$_{59}$N$_2$O$_4$ | Newly synthesized | NA | ND | |
| 372.3113 | 19.96    | C$_{21}$H$_{31}$NO$_4$ | Newly synthesized | T. versicolor and G. applanatum | 3.6E+05 | +
| 377.2302 | 20.09    | C$_{19}$H$_{32}$N$_2$O$_3$ | Newly synthesized | NA | ND | |
| 400.1019 | 7.96     | C$_{17}$H$_{33}$N$_2$O$_2$ | Newly synthesized | NA | ND | |
| 432.1504 | 8.88     | C$_{19}$H$_{31}$N$_2$O$_7$ | Newly synthesized | NA | ND | |
| 474.3068 | 15.86    | C$_{24}$H$_{43}$NO$_4$ | Newly synthesized | T. versicolor | 1.25E+04 | +
| 492.3176 | 15.04    | C$_{24}$H$_{43}$NO$_4$ | Newly synthesized | T. versicolor | 1.50E+04 | +
| 496.2886 | 16.43    | C$_{15}$H$_{41}$N$_2$O$_4$ | Newly synthesized | T. versicolor | 4.0E+04 | +
| 506.3332 | 15.85    | C$_{25}$H$_{47}$NO$_4$ | Newly synthesized | T. versicolor | 3.0E+04 | +
| 599.2043 | 8.63     | C$_{31}$H$_{54}$O$_{12}$ | Newly synthesized | NA | ND | |
| 716.5259 | 18.91    | C$_{42}$H$_{68}$NO$_3$ | Newly synthesized | NA | ND | |

Yellow label, the features were observed by LC-MS in both negative and positive modes. These features were only counted once. Plus mark, the features discovered in this work were tagged with plus mark. NA, the features had no $^{13}$C incorporation. ND, the features had weak MS signal intensity or no signal in the supernatants from the co-culture on day 10. Fold increase, fold value was the ratio of MS signal intensity of increased feature in the co-culture to that in the control mono-culture. Data show the mean with standard deviation calculated from three independent biological replicates.
were added, only 20 mL of fresh medium with $^{13}$C-labeled glucose as carbon source. After the mono-culture for 10 or 20 days, 10 mL of \textit{T. versicolor} and \textit{G. applanatum} culture broths were harvested for analysis of $^{13}$C-labeling. All samples had three independent biological replicates and two analytical replicates. Samples were extracted and analyzed by LC-QTOF-MS as described above. The mass isotopomer distributions were corrected for the contribution from natural isotopes by a matrix-based method (Jennings and Matthews, 2005). The total $^{13}$C-incorporation for each feature was obtained by normalizing its total carbon number as in our previous publication (Yang et al., 2013). Relative isotopic abundance ($M_i$) for a feature in which $i$ $^{13}$C atoms were incorporated was calculated by the Equation (1):

$$M_i(\%) = \frac{m_i}{\sum_{j=0}^{n} m_j}$$  

where $m_i$ represents the isotopic abundance for a feature in which $i$ $^{13}$C atoms were incorporated and $n$ represents the maximum number of $^{13}$C atoms incorporated.

Total $^{13}$C-incorporation of a feature with $N$ carbon atoms was obtained by normalizing to its total carbon number according to the Equation (2):

$$\text{Total }^{13}\text{C} - \text{incorporation (\%) } = \frac{\sum_{i=1}^{N} i \times M_i}{N}$$  

The significance of difference of $^{13}$C-incorporation between experimental data points was determined by $t$-tests (Origin 8.0). A $P$-value <0.05 was considered to be statistically significant.

**Isolation and Purification of Compound 1**

Twenty liters of the co-cultured supernatant was extracted three times with ethyl acetate (EtOAc). 2.5 g of crude EtOAc extract was concentrated under reduced pressure and purified on a silica gel (200–300 mesh) column and then eluted with petroleum ether/ethyl acetate and chloroform/methanol system to yield ten fractions (Song et al., 2014). Eight hundred and fifty milligrams of fraction 6 and 7 from the chloroform/methanol elution (20:1 and 10:1, v/v) was used to a medium pressure liquid chromatography (Flash CO140080-0, Agela Technologies, China) and eluted with methanol/water (the elution gradient was 10–90% methanol in 65 min) to generate a mixture of 216 mg. This mixture was purified on a silica gel (200–300 mesh, Qingdao Haiyang Chem. Ind. Co. Ltd. China) column by eluting with petroleum ether/ethyl acetate/methanol (3:3:1, v/v) and then separated on a preparative column (Venusil XBP C18, Agela Technologies, China) by eluting with methanol/water (the elution gradient was 10–50% methanol in 40 min and flow rate was 8 mL/min) to obtain 10 mg of compound 1.

**NMR Analysis of Purified Compound 1**

$^1$H, $^{13}$C, and 2D NMR spectra of the purified compound 1 were all performed by using a Bruker Avance 600 MHz spectrometer (Karlsruhe, Germany) at 25°C. The compound 1 was accurately weighted, and dissolved in 0.5 mL of deuterated methanol as the internal lock. The resulting spectra were manually phased and baseline corrected and calibrated to methanol, using TOPSPIN (Version 2.1, Bruker).

**Cell Viability Assay**

The human leukemic cell line U937 and lung cancer cell line A549 were purchased from American Type Culture Collection (Manassas, VA, USA). They were cultured in RPMI-1640 (HyClone, USA) with 10% FBS (fetal bovine serum, Gibco). Cells were inoculated in 96-well plates at a density of 3,000 cells per well overnight and then treated to different concentrations of compound 1 for 48 h. The compound 1 was solved in ethanol. At a concentration of 300 \(\mu\text{M}\) of compound 1, the ethanol was 0.8\% (v/v) in the cell culture medium, showing minimal effect on the viability of both cell lines. The effect of compound 1 on the viability of U937 and A549 cells was evaluated with CellTiter 96® AQueous One Solution (Promega, Madison, WI, USA) (Soman et al., 2009). The absorbance was read at 490 nm with a microplate spectrophotometer (Multiskan FC, Thermo scientific, USA). Three independent experiments were performed and each one had six replicates. Data show the mean with error bar indicating standard deviation calculated from biological replicates by Origin 8.0. IC$_{50}$ defined as the concentration with the inhibition of 50% cells was calculated by using SPSS (Statistical Package for Social Sciences) package 6, version 15.0.

**Antioxidant Activity Assay**

The 2,2′-azinobis-(3-ethylbenzothiazoline-6-sulfonic acid) (ABTS) method used is based on the reduction of the ABTS$^{+}$ radical action by the antioxidants present in the sample (Rigano et al., 2014). Three biological replicates were performed. A solution of 7.4 mM ABTS$^{+}$ (5 mL) mixed with 140 mM K$_2$S$_2$O$_8$ (88 \(\mu\text{L}\)) was prepared and stabilized for 12 h at 4°C in the dark. This mixture was then diluted by mixing ABTS$^{+}$ solution with ethanol (1:88) to obtain an absorbance of 0.70 ± 0.10 unit at 734 nm using a spectrophotometer (Multiskan FC, Thermo scientific, USA). Compound 1 (100 \(\mu\text{L}\)) and the standard controls (ascorbic acid and phenolic antioxidant 2,6-di-tert-butyl-4-methoxyphenol) reacted with 1 mL of diluted ABTS$^{+}$ solution for 2.5 min, and then the absorbance was taken at 734 nm against a blank constituted by ABTS$^{+}$ solution added with 100 \(\mu\text{L}\) of ethanol. ABTS$^{+}$ scavenging activity was calculated by the Equation (3):

$$\text{Scavenging effect (\%)} = \left(1 - \frac{\text{Abs. sample}}{\text{Abs. blank}}\right) \times 100$$  

where Abs blank = 100 \(\mu\text{L}\) of ethanol+1 mL of diluted ABTS$^{+}$ solution.

**RESULTS**

**Induced Feature Discovery in Fungal Interaction**

Unsupervised PCA is well-suited for comparing different biological samples and identifying statistically significant differences (Chen et al., 2007). As shown in Figure 1 (left figure), examination of the scores plot showed that the co-culture treatment is clearly separated from the two mono-cultures. The variable features responsible for discriminating these three groups are shown in the loading plots (Figure 1, right). More
than 4,000 features were recorded, out of which 58 features were detected only in the co-culture and 16 features were at least 5-fold more abundant in the co-culture than those in the control mono-culture (Table 1, Figure 2). Compared to our previous work (Yao et al., 2016), 28 additional features were discovered mainly due to the optimized chromatography (Table 1, Figure 2).

**13C-Labeling Analysis to Associate Induced Features with Corresponding Fungus**

To associate induced features with corresponding fungus, we initially harvested individual mycelium pellets of *T. versicolor* and *G. applanatum* after the co-culture for 10 days and then detected the abundance of *in vivo* induced features. Many of the induced features were observed for both samples of *T. versicolor* and *G. applanatum* (data not shown). This made it impossible to distinguish if individual or both fungi were induced to generate the compounds in the co-culture. To overcome this issue, we designed a 13C-labeling approach in the liquid co-cultivation. The workflow is shown in Figure 3. First, *T. versicolor* and *G. applanatum* were co-cultivated for 10 days to activate the cryptic genes. Next, their mycelia were steriley separated and mono-cultivated in the liquid medium which contained half of fresh medium with 13C-labeled glucose as carbon source and half of co-cultured supernatants collected on day 10. Then, the samples were harvested in the mono-cultures of *T. versicolor* and *G. applanatum* on days 10 and 20 and analyzed by LC-QTOF-MS. In the preliminary experiments, the samples were also harvested after 5 days of mono-culture with the addition of the supernatants. However, many of induced features were only slightly labeled. It was likely due to a relatively long lag phase and low growth rate in the mono-culture of *T. versicolor* and *G. applanatum* after the stimulation of the co-culture. In addition, for the unlabeled features, 13C-labeling was not detected on day 5 either. Therefore, the samples were harvested later in order to obtain the strong MS signal. As the incorporation of 13C-labeled carbons from glucose increases the molecular weight of metabolites, the mass shift determined from the mass spectra...
FIGURE 2 | Overview of the number of the induced features in the co-culture of T. versicolor and G. applanatum.

FIGURE 3 | $^{13}$C-labeling analysis procedure. T. versicolor and G. applanatum were co-cultivated for 10 days, and then their mycelia were sterilely separated and mono-cultivated in the liquid medium which contained half of fresh medium with $^{13}$C-labeled glucose as carbon source and half of co-cultured supernatants. Then samples were harvested in the mono-cultures of T. versicolor and G. applanatum and analyzed on LC-QTOF-MS.

provides then the information about which fungi generated the induced features.

Total 31 induced features were found to have $^{13}$C incorporation in the mono-culture, and 43 features did not incorporate any label (Figure 2). Among the labeled features activated by the co-culture, 20 originated from T. versicolor and 6 features were derived from G. applanatum. Five features were induced by both T. versicolor and G. applanatum (Figure 2). Several representative $^{13}$C labeling results are shown in Figure 4. In Figure 4A, the m/z 490 (experimental m/z 490.3025, predicted elemental component $C_{24}H_{45}NO_9$, error 0.3 ppm) was a newly synthesized feature in the co-culture. As $^{13}$C-labeled carbon was incorporated into the feature m/z 490, intensity of m/z 491 ($m/z +1$) up to m/z 506 ($m/z +16$) increased. This change
A comparison of relative MS intensity suggests that the induced features have $^{13}$C incorporation with time. (A–D) Upper figures: representative mass spectral data for 10 day time point showing that mass intensity caused by $^{13}$C incorporation in the mono-culture of *T. versicolor* or *G. applanatum*. Lower figures: comparison of total $^{13}$C incorporation shows that induced feature was produced by either *T. versicolor* or *G. applanatum*, or both. Data show the mean with error bars indicating standard deviation calculated from three independent replicates (*P* ≤ 0.05, t-test). (E) There was no $^{13}$C incorporation for N-(4-methoxyphenyl)formamide 2-O-beta-D-xyloside on day 10 and 20. was observed only in the mono-culture of *T. versicolor* and indicates that m/z 490 was particularly induced in this species. The calculation of total $^{13}$C incorporation indicated that about 19 and 25% carbon was replaced with $^{13}$C on days 10 and 20, respectively (bottom part of Figure 4A). The m/z 167.0348 in Figure 4B was highly produced during the co-culture and identified as orsellinic acid in the previous publication (Yao et al., 2016). In contrast to the above example, the m/z 167 incorporated $^{13}$C-labels in the same time frame only in the mono-culture of *G. applanatum*, resulting in the parent ions shifted from m/z 167 to m/z 174 in the $^{13}$C-labeled mass spectra and total $^{13}$C incorporation level reached 14% on day 20 (Figure 4B). The m/z 165.0554 was previously identified as 3-phenyllactic acid (Yao et al., 2016), and $^{13}$C-labeled mass spectra ranging from m/z 165 to m/z 174 were observed in both mono-cultures of *T. versicolor* and *G. applanatum* (Figure 4C). Similar example, feature with m/z 309 (experimental m/z 309.0756, $C_{18}H_{14}O_5$, error 0.37 ppm) increased 15.4-fold during the co-culture in comparison with MS signal in the control mono-culture and $^{13}$C-labeling was observed for both fungi (Figure 4D). It is worth mentioning that a contamination from the undesired fungus cannot be absolutely excluded during the transfer from the co-culture to the mono-culture, but that it does not affect the identification of the fungus producing a feature only found in the mono-culture of either *T. versicolor* or *G. applanatum* based on the $^{13}$C-labeling analysis.

$^{13}$C incorporation could be due to the induction of features which have been released into the medium during the co-culture. To confirm whether diffusible features were involved in triggering the silent gene expression in this study, we treated *T. versicolor* and *G. applanatum* with fresh medium containing $^{13}$C labeled glucose as carbon source but without the addition of the supernatant of co-culture on day 10. As shown in Figure 5 as an example, $^{13}$C labeling of m/z 490.3025 and 136.0403
was not observed up to 20 days but when the supernatant was added the features were significantly labeled. Notably, for 12 induced features with high signal intensities, including newly synthesized xyloside [N-(4-methoxyphenyl)formamide 2-O-beta-D-xyloside; Yao et al., 2016], we did not detected any $^{13}$C incorporation in the mono-culture of either fungus (Table 1, Figure 1). Therefore, it was not possible to assign their origin to a specific fungus.

**Dereplication of Newly Discovered Features by Molecular Network Analysis**

To investigate the structural similarities occurring for 28 newly discovered features in this study, MS/MS fragmentation spectra of the induced features in Table 1 were processed and organized as molecular network with the previously identified features. Figure 6 (upper part) shows a constructed subnetwork to dereplicate m/z 196.0944 and 216.1021. These two new features likely possessed similar backbone structure with the previously identified N-(2-hydroxy-4-methoxyphenyl)formamide (m/z 168.0653) and its analogous features (m/z 140.0708, 150.0548, 230.1177, 287.1030) due to their close fragmentation patterns (Supplementary Figure 2). This data was also in agreement with $^{13}$C-labeling analysis, in which most of features involved in this subnetwork were not labeled either.

In another example, five newly identified features and three previous features (m/z 251.0712, 281.0808, and 334.0733) were clustered together with high scores (Figure 6, bottom part). Comparison of the fragment ions of m/z 306.0775, 334.0733, 581.1208, 629.1419, and 279.0656 showed some common ions of m/z 77.04, 117.03, 235.08 which were likely derived from fragmentation of m/z 279.0656, suggesting that these features probably had the same backbone structure and belonged to a series of structural derivatives (Supplementary Figure 2). Moreover, $^{13}$C incorporation of m/z 279.0656, 251.0712, and 281.0808 were clearly detected in the mono-culture of *T. versicolor* with the addition of the supernatant of co-culture (Supplementary Figure 1), suggesting that the derivatives of m/z 279.0656 (i.e., m/z 306.0775, 334.0733, 581.1208, and 629.1419), which did not show $^{13}$C incorporation due to the weak signals in the mono-culture) were also likely biosynthesized by *T. versicolor* under the co-cultured condition.

**Identification of Induced Compound 1 and Analysis of Its Biological Activity**

Since compound 1 was derived from both *T. versicolor* and *G. annaplanum* (Figure 4D), and 4.4-fold more abundant in the co-culture, we isolated and purified sufficient amount for detailed characterization. Compound 1 was a pale yellow powder with the molecular formula of C$_{18}$H$_{14}$O$_{5}$. The $^{13}$C NMR and HSQC spectrum showed the presence of two carbonyl carbon, one oxygen connected CH, one oxygen connected to quaternary carbon, two single benzene rings and two olefinic carbon atoms (one with oxygen attached to it) (Supplementary Table 1 and Supplementary Figure 3). All the information suggested that the basic skeleton of the compound 1 is a terphenyl derivative with two mono-substituted benzene rings. The COSY spectrum showed the $^{1}$H–$^{1}$H spin systems of H-2'/H-3'/H-4'/H-5'/H-6' and H-2'/H-3'/H-4'/H-5'/H-6'' assigned two mono-substituted benzene rings (A and B). The HMBC correlations from H-2 ($\delta_{H}$ 4.49) to C1 ($\delta_{C}$ 203.4), C3 ($\delta_{C}$ 197.4), C4 ($\delta_{C}$ 113.5), and C6 ($\delta_{C}$ 90.9), and from OH-6 ($\delta_{H}$ 7.98) to the carbonyl C1 ($\delta_{C}$ 203.4). The linkage of the ring B to the ring C at C-4 was confirmed by HMBC correlations from H-2'' ($\delta_{H}$ 7.88) to C3 ($\delta_{C}$ 113.5) and C-1''($\delta_{C}$ 135.4). Therefore, the structure of compound 1 was identified as a phenyl polyketide, 2,5,6-trihydroxy-4,6-diphenylcyclohex-4-ene-1,3-dione (Figure 7).

We further tested biological activity of compound 1. The human lung cancer cell lines A549 and leukemic cell lines U937 were treated with compound 1 at various concentrations for 48 h. As shown in Figure 8A, no visible changes in cell viability were detected for human lung cancer cell line A549 when the concentrations were increased to 300 $\mu$M. In contrast, compound 1 inhibited the viability of leukemic cells in a dose-dependent manner. The IC$_{50}$ at 48 h was determined
to be $276 \pm 5\, \mu M$ (equal to 85 mg/L). In addition, based on the structural characteristics of compound 1, we also studied whether compound 1 had antioxidant properties using ABTS assay (Figure 8B). The highest percentage of antioxidant capacities ($82.65 \pm 1.25\%$) was observed for compound 1 at the concentration of $200\, \mu g/mL$. This is comparable with the report of crude extracts from berries (Abu-Bakar et al., 2016). The comparison of the antioxidant activity with ascorbic acid (VC) and phenolic antioxidant 2,6-di-tert-butyl-4-methoxyphenol (BHT) revealed that ascorbic acid $>$ compound 1 $>$ BHT.

**DISCUSSION**

Determining the origin of induced features in the co-culture remains a large challenge, because it generally requires the chemical structures with their biological information. Previously Bertrand et al. discovered five de novo induced compounds from the co-culture of *Trichophyton rubrum* and *Bionectria ochroleuca*, and elucidated the origin of one of them based on its non-sulfated form detected in the mono-culture of *B. ochroleuca* (Bertrand et al., 2013). However, other four compounds could not be associated with corresponding fungi due to the lack of their structures. In another study Ola et al. isolated nine compounds and speculated that four of them detected only in the co-culture originated from *Fusarium tricinctum* in terms of structural analogies with the known fungal products from the Xylariaceae family, but the producer of remaining five compounds was still unclear because of insufficient biochemical evidences (Ola et al., 2013). In this work, among 74 induced features, 31 features were shown to be produced specifically by either *T. versicolor* or *G. applanatum*, or by both fungi using a $^{13}$C-based labeling analysis. This methodology was able to distinguish the origin even if the identities of compounds were not available or almost nothing was known about biochemical aspects. In more detail, the total $^{13}$C incorporation in the same fungal culture varied noticeably among features. Isotopic steady state is the state that $^{13}$C-labeling signatures in metabolites become time invariant (Antoniewicz et al., 2007). In current
work, $^{13}$C incorporation into 20 features (e.g., orsellinic acid) had the similar abundance levels between days 10 and 20 and these features had the similar $^{13}$C-labeling pattern (data not shown), indicating that they likely reached isotopic steady state around 10 days (Figure 4B, Supplementary Figure 1). In contrast, $^{13}$C incorporation level of 11 features including novel phenyl polyketide (compound 1) increased from day 10 to day 20 (Figures 4A,C,D, Supplementary Figure 1), suggesting either the fluxes via their synthetic pathways were low or their metabolite pools were relatively large (Zamboni et al., 2009).

By comparison of $^{13}$C-labeling patterns between with and without the addition of the supernatant of co-culture, it was demonstrated that the induction of gene expression and synthesis of corresponding metabolites during the co-culture of T. versicolor and G. applanatum depended indeed on the inter-mycelial communication. Forty-three induced features in the supernatant of co-culture on day 10 were detected with signal intensity ranging from $10^3$ to $10^5$ (Table 1), but in the current work we were not able to determine which features were responsible for signaling to activate gene expression in T. versicolor or G. applanatum. Additional studies will be needed to confirm this link as well as to elucidate the structure and function of signaling molecules. Notably, the previous report also demonstrated that an intimate physical interaction of the actinomycete S. hygroscopicus and fungal mycelia of Aspergillus nidulans was required to induce specific stimulation of the silent polyketide synthases and non-ribosomal peptide synthetases gene clusters (Schroechk et al., 2009). In our research, we found 12 highly produced features that did not incorporate $^{13}$C labeling in the mono-culture. One potential explanation is that these features were also produced in the co-culture via mycelium physical interaction to elicit the specific response. In addition, compound 1 had over 15-fold higher MS signal intensity in the co-culture than in the control mono-culture of T. versicolor where the mycelia were not pre-induced by the co-culture. It was not detected in the control mono-culture of G. applanatum, yet $^{13}$C incorporation in G. applanatum was 1.3-fold higher than that in T. versicolor on day 20 (Figure 4D). This finding suggests that the co-culture could significantly activate the encoding genes or gene clusters related to the synthesis of Compound 1 in G. applanatum. This provides an important insight into possible manipulation of G. applanatum to enhance biosynthesis of novel metabolites.

Comparing MS fragmentation similarity including common losses, molecular network analysis is able to obtain a simultaneous visual investigation of identical molecules, analogs, or metabolite families, thereby assisting the structural analysis (Winnikoff et al., 2014; Cabral et al., 2016). In our previous research, this method was utilized to show that the common neutral loss of 132 Dalton resulted from the deglycosylation reaction, which helped to find a series of novel xylosides generated during the co-culture (Yao et al., 2016). Here, combined molecular network analysis and $^{13}$C-labeling analysis suggested further that some newly discovered features were not only structurally analogous but also had similar induction mechanism and were likely produced by the same fungus. Thus, combination of network analysis and $^{13}$C labeling shows promise to accelerate the elucidation of biosynthetic pathways of novel metabolites.

Several type II polyketides have been reported to be high-value medicinals (Zhang W. et al., 2012). In antioxidant assay, compound 1 had better activity than BHT. More interestingly, compound 1 at micromolar concentration was able to inhibit the viability of leukaemic cells. For comparison, Zhang et al. reported that matrine extracted from Sophora flavescens inhibited the proliferation of acute myeloid leukemia cell lines U937 in a dose- and time-dependent manner with the IC$_{50}$ of 590 mg/L at 24 h, and resulted in the maximal apoptosis rates with 37.2% for 24 h (Zhang S. et al., 2012). Wang et al. also demonstrated that 100 $\mu$g/mL of Ganoderma lucidum extracts could greatly suppress leukaemic cell growth with the inhibition rate of 75% (Wang et al., 1997). In our case, the IC$_{50}$ of compound 1 was at the same level with matrine and crude extracts from G. lucidum. Thus, it can become a starting point for development of lead compounds to cure leukemia or other cancers.

**CONCLUSION**

The application of $^{13}$C-labeling analysis produced valuable insight into the role of individual partners in the co-culture in production of known or unknown induced metabolites. Moreover, $^{13}$C-labeling approach combined with molecular network analysis can reveal whether certain structural analogs were produced by the same fungus or through the similar activation mechanism. The current $^{13}$C-labeling information sets an important foundation for further studies in the basidiomycetes, including but not limited to novel metabolites discovery and biosynthetic capacity improvement.
FIGURE 8 | The biological activity study of compound 1. (A) Addition of compound 1 affected the cell viability of U937 but not of A549. (B) Graph of ABTS•+ scavenging activity vs. concentration. Data show the mean with error bars indicating standard deviation calculated from three independent biological replicates. VC, ascorbic acid; BHT, 2,6-di-tert-butyl-4-methoxyphenol.

AUTHOR CONTRIBUTIONS

X-YX, BQ, and SY: conceived and designed the project; X-YX, X-TS, X-JY, Y-MZ, HF, and JY: performed the experiments; X-YX, X-TS, L-PZ, F-YD, MS, and SY: interpreted the data. All authors contributed to the preparation of the manuscript, read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

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