A new species of *Trichoderma hypoxylon* harbours abundant secondary metabolites

Jingzu Sun1,2, Yunfei Pei1,†, Erwei Li1, Wei Li1, Kevin D. Hyde2, Wen-Bing Yin1,3 & Xingzhong Liu1,3

Some species of *Trichoderma* are fungicolous on fungi and have been extensively studied and commercialized as biocontrol agents. Multigene analyses coupled with morphology, resulted in the discovery of *T. hypoxylon* sp. nov., which was isolated from surface of the stroma of *Hypoxylon anthochroum*. The new taxon produces *Trichoderma*- to *Verticillium*-like conidiophores and hyaline conidia. Phylogenetic analyses based on combined ITS, TEF1-α and RPB2 sequence data indicated that *T. hypoxylon* is a well-distinguished species with strong bootstrap support in the polysporum group. Chemical assessment of this species reveals a richness of secondary metabolites with trichothecenes and epipolythiodiketopiperazines as the major compounds. The fungicolous life style of *T. hypoxylon* and the production of abundant metabolites are indicative of the important ecological roles of this species in nature.

Traditionally the taxonomy of species of *Trichoderma* was based on morphology. Most species in this genus are usually fast growing, produce highly branched conidiophores with cylindrical to nearly subglobose phialides and ellipsoidal to globose conidia1–4. However, high morphological homoplasy in sexual state makes identification difficult, and the importance of sequence data have been increased5–6. Based on the combined phenotypic and phylogenetic analysis, about 260 species have been recognized and accepted5–10. The internal transcript spacers (ITS), translation elongation factor 1-alpha (TEF1-α) and largest subunit of RNA polymerase II (RBP2) genes are more available to recognize species within *Trichoderma*5,9,10. Phylogenetic analysis based on a combination of ITS, TEF1-α and RPB2 are recommended solve the problem of *Trichoderma* species complex and reveal taxonomic diversity2,6–11.

Fungi associated with other fungi as saprobes, commensals or parasites are termed fungicolous fungi12. These fungi usually produce rich secondary metabolites, which have been reported as an important resource for bioactive small molecule discovery such as anti-fungi, tumors, nematodes and bacteria13,14. Some metabolites produced by the species in this genus play very important ecological roles in nature15. *Trichoderma* species are most frequently found in vegetable matter, decaying wood and soil, plant rhizosphere, as well as on other fungi5,6,16. Fungicolous *Trichoderma* species comprise diversity and abundant genes associated with secondary metabolites productivity15,16. These genes are responsible to a number of secondary metabolites with pharmaceutical and biotechnological importance including peptides, peptaibols, poliketides, pyrones, siderophores and nonvolatile terpenes14,15,21–22. Some of secondary metabolites are responsible for survival and adaption of their habitat15,16. For example, peptaibols are small peptides of non-ribosomal origin produced by *Trichoderma*15 and have the ability to induce systemic resistance in plants against microbial invasion. Harzianum A is a growth-promoting trichothecene produced by *Trichoderma arundinaceum*17.

In the search of novel bioactive compounds, we carried out resource and diversity investigation of fungicolous fungi in China and Thailand. Two isolates of *Trichoderma* were obtained from stroma of *Hypoxylon anthochroum*. They are described, illustrated and named as a new species *Trichoderma hypoxylon*. Its phylogenetic positions

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1State Key Laboratory of Mycology, Institute of Microbiology, Chinese Academy of Sciences (CASMIM), No. 3 Park 1, West Beichen Road, Chaoyang District, Beijing 100101, China. 2Center of Excellence in Fungal Research, and School of Science, Mae Fah Luang University, Chiang Rai, 57100, Thailand. 3Joint Laboratory of Applied Microbial Technology, CASIM and Institute of Biology limited Liability Company, Henan Academy of Sciences, Zheng Zhou, 45002, China. †Present address: National Institutes for Food and Drug Control, No. 2, Tiantan Xili, Dongcheng District, Beijing 100050, China. Correspondence and requests for materials should be addressed to W.-B.Y. (email: yinwb@im.ac.cn) or X.L. (email: liuxz@im.ac.cn)
were also explored, which inferred from sequence analyses of the combined internal transcribed spacer (ITS), partial RNA polymerase II subunit (RPB2) and translation elongation factor 1 alpha (TEF1-α) exon genes.

**Results**

**Phylogenetic analyses.** Alignment results show that the sequences ITS, RPB2 and TEF1-α of *T. hypoxylon* are less than 97% similar to other *Trichoderma* species. Based on phylogenetic analysis of single gene of ITS, RPB2 and TEF1-α, *T. hypoxylon* formed a clade with *Trichoderma taxi* and *Trichoderma rubi*. The position of this clade showed closed relationship with section Hypoceanum, Polysporum, Psychrophila and other species in *Trichoderma* (Fig. S1, and Fig. S2). The phylogenetic analysis of RPB2 showed that *T. hypoxylon* grouped with the Polysporum section (Fig. S3). Therefore, the combination matrix included 60 ingroup taxa of *Trichoderma* which are phylogenetically close to *T. hypoxylon*. This data matrix comprised 2839 characters. Bayesian inference (BI), maximum likelihood (ML) and maximum parsimony (MP) trees generated shared the same topology. In MP analyses, 1642 (57.8%) characters are constant, 216 (7.6%) characters are parsimony-uninformative, and 981 (34.6%) characters are parsimony-informative. In MP analyses, 1642 characters were constant, 981 were parsimony-informative, and 216 variable characters were parsimony-uninformative. Maximum likelihood tree was presented (Fig. 1).

Based on the analyses, 60 strains of *Trichoderma*, including our new species *T. hypoxylon*, formed a strongly supported group (MPBP/MLBP/BIPP = 100%/100%/1.00). They also clustered in 7 recognized subclades and a new subclade including *T. hypoxylon*, Brevicompactum (MLBP/BIPP = 94%/1.00), Deliquescent (MPBP/BIPP = 100%/100%/1.00), Hypoceanum (MPBP/MLBP/BIPP = 100%/99%/1.00), Polysporum (BIPP = 1.00), Psychrophila (MPBP/MLBP/BIPP = 98%/96%/1.00), and green spore sections (MPBP/MLBP/BIPP = 100%/100%/1.00), whereas two strain of *Trichoderma atroviride* Bissett positioned inside Polysporum section. The tree topology is basically congruent with previous reports6.
The new species *T. hypoxylon* (CGMCC 3.17906, CGMCC 3.17907), and *T. taxi* and *T. rubi* formed a clade independent from other *Trichoderma* species with strong support (MPBP/MLBP/BIPP = 100%/100%/1.00). However, *T. hypoxylon* isolates were distinguished from *T. taxi* and *T. rubi* with high support (MPBP/MLBP/BIPP = 100%/100%/1.00).

Figure 2. Morphological characteristics of *Trichoderma hypoxylon* (ex-type CGMCC 3.17906) on PDA. (a–c) and (h–k) conidiophores with conidia; (d–g) conidiophores without conidia; l, conidia; bar = 10 μm.
Figure 3. Morphological characters of *Trichoderma hypoxylon* (ex-type CGMCC 3.17906) on CMD after 30 days. (a) Forward of the colony; (b) reverse of the colony; (c) mycelia; (d,e,i) conidiophores with conidia; (h) conidiophores without conidia; (j) conidia; c = 1000 μm; (d) e = 20 μm; c f-j = 10 μm.
Taxonomy. *Trichoderma hypoxylon* Jing Z. Sun, Xing Z. Liu & K.D. Hyde, *sp. nov.*

**Index Fungorum number:** IF552046, **Facesoffungi number:** FoF: 02075, Figs 2–4.

**Type:**—HMAS 246918

Colonizing or hyperparasitic on stroma of *Hypoxylon anthochroum*. Sexual morph Undetermined. Asexual morph forming *Acremonium-* to *Verticillium*-like conidiophores and hyaline conidia. Phialides lageniform to cylindrical, hyaline, 4.5–12 × 3–3.5 μm (\(\bar{x} = 6.5 \pm 1.4 \times 3.2 \pm 0.25 \mu m, n = 30\)); l/w 2.5–4 (−4.5) (n = 30). Mature conidia forming a head at the apex of conidiophores, hyaline, smooth-walled, unicellular, oboviod and mostly 2.8–4.3 × 2.1–2.4 μm (\(\bar{x} = 3.8 \pm 0.4 \times 2.2 \pm 0.10 \mu m, n = 30\)), l/w 1–2 (−2.2) (n = 30). Slow growing on PDA, SNA and CMD; PDA colonies especially dense, whitish; SNA and CMA colonies consisting of concentric rings with irregular outline when cultured in darkness at 30 °C.

Colonies on PDA after 10 d at 20, 25 and 30 °C dense (Fig. 4a–c), with a thick white layer of cotton-like aerial mycelia, forming concentric rings at 20 and 25 °C, however not forming concentric rings at 30 °C, odour woody, and agar not pigmented. Conidia formed within 10 d in the aerial mycelium, mature conidia gathered at the apex of conidiophores, hyaline, smooth walled, 1-celled, oboviod and mostly 2.8–4.3 × 2.1–2.4 μm (\(\bar{x} = 3.8 \pm 0.4 \times 2.2 \pm 0.10 \mu m, n = 30\)), l/w 1.2–2.2 (−2.5) (n = 30).

Colonies on CMD after 10 d at 20, 25 and 30 °C flat (Fig. 4d–f), with a thin white layer of mycelia, not forming concentric rings and conidia at 20 and 25 °C, however forming concentric rings at 30 °C, no distinctive odour, agar not pigmented. Conidiophores *Acremonium-* to irregularly *Verticillium*-like; phialides lageniform to cylindrical, hyaline, 8.5–19.5 × 2–2.51 μm (\(\bar{x} = 13.5 \pm 1.4 \times 2.1 \pm 0.18 \mu m, n = 30\)), l/w 3.5–9 (−9.5) (n = 30); Conidia formed after 20 d, mature conidia gathered at the apex of conidiophores hyaline, smooth walled, one-celled, oval, 4.8–6.3 × 3.0–3.4 μm (\(\bar{x} = 4.9 \pm 0.4 \times 3.1 \pm 0.1 \mu m, n = 30\)), l/w 1.2–1.6 (−2.0) (n = 30).

Figure 4. Cultures of *Trichoderma hypoxylon* on PDA, CMD and SNA at different temperature after 10 days. (a–c) Colony on PDA at 30 °C, 25 °C and 20 °C seperately; (d–f) colony on CMD at 30 °C, 25 °C and 20 °C seperately; (g–i) colony on SNA at 30 °C, 25 °C and 20 °C seperately.
Colony on SNA after 10 d at 20, 25 and 30 °C flat (Fig. 4g–i), with a thin white layer of mycelia, not forming concentric rings and conidia at 20 and 25 °C, however forming concentric rings at 30 °C, no distinctive odor, agar not pigmented.

Colony radius on PDA after 72 h at 20 °C, 28.2–30.9 mm; 25 °C, 33.4–36.3 mm; 30 °C, 0.32–1 mm; and 35 °C, 0 mm (n = 5). Colony radius on CMD after 72 h at 20 °C, 41.2–46.1 mm; 25 °C, 37.4–38.3 mm; 30 °C, 8.32–9.12 mm; and 35 °C 0 mm (n = 5). Colony radius on SNA after 72 h at 20 °C, 24.9–26.8 mm; 25 °C, 27.1–29.2 mm; 30 °C, 10.4–12.1 mm, and 35 °C, 0 mm (n = 5).

Etymology:—hypoxylon refers to the genus of host fungus *Hypoxylon anthochroum*.

Distribution:—Chiang Rai, Thailand

Host:—*Hypoxylon anthochroum*, a saprobic fungus on dead wood.

Material examination:—Thailand, Chiang Rai Province, isolated from the stroma of *Hypoxylon anthochroum*, 5 May, 2014. MFLU16–1263 (Holotype, dried culture MFLU 16–1263!), HMAS 246918 (Isotype, dried culture HMAS 246918) ex-type living culture, MFUCC 15–0683, CGMCC 3.17906, CGMCC 3.17907.

Notes: The asexual morph of *Trichoderma hypoxylon* produces *Acremonium*- to *Verticillium*-like conidio- phores and hyaline conidia. Phylogenetically *T. hypoxylon* is related to *T. taxi* and *T. rubi* and together formed a new independent clade distinguished other *Trichoderma* species (Fig. 1). However, *T. hypoxylon* can be phylogenetically distinguished from these species. The conidia of *T. hypoxylon* (2.8–4.3 × 2.1–2.4 μm) are also longer than *T. taxi* (2.4–3.1 × 2.0–2.5 μm)23 and *T. rubi* (2.3–3.3 × 2.0–2.7 μm)6. Meanwhile, *T. hypoxylon* did not produce pigment on medium, whereas *T. rubi* produced brownish pigment and yellow crystals6.

Secondary metabolite analysis and characterization of compounds. HPLC analyses of extracts of *T. hypoxylon* cultivated on PDA medium were carried out to assess the production of secondary metabolites. Four major peaks with high yields were obtained by HPLC chromatography (Fig. 5A). To characterize compounds from *T. hypoxylon*, 1 liter fermentation on PDA medium were performed. After the semi-preparative reversed-phase HPLC separation step, we isolated four known compounds trichodermamide A (1), aspergillazine A (2), aspergillazine C (3) and harzianum B (4) (Fig. 5B). The assignments of four compounds were based on the published data of proton NMR24–27. Notably, the yields of harzianum B were 50 mg out of 330 mg crude extracts.

Discussion

Traditionally, delimitation of *Trichoderma* species was mainly based on the morphology1, although, it could not well explain the taxonomic position of these species6–9. Phylogenetic analysis has resulted in the discovery of many new species and has been extensively used in fungal taxonomy28. Presently, more than 258 species of *Trichoderma* are accepted based on phylogenetic analysis5,19. However, many species of *Trichoderma* remain to be discovered and described5,6. To explore the taxonomic position of *T. hypoxylon*, phylogenetic tree containing all species of *Trichoderma* species was constructed and the putative position of *T. hypoxylon* was shown (Figs S1–S3). The single gene of ITS, RPB2 and TEF1-α could distinguish *T. hypoxylon* from other *Trichoderma* species, the results suggested that these three genes are effective in taxonomy of *Trichoderma*5,6,9. However, the single gene could not well delimit Longibrachiatum, Viride and some other sections in genus of *Trichoderma*. Multigene analysis was a popular and feasible approach to solve the problem5,6,10.

Based on analyses of the combined sequences of ITS, RPB2 and TEF1-α, 56 currently known species in *Trichoderma* clustered together (Fig. 1). Nine subclades, Brevicompactum, Deliquesceans, Green spore group,
Hypoenceum, Longibrachiatum, Polysporum, Psychrophilium, Viride and a new subclade including *Trichoderma hypoxylon* were recognized, which is basically congruent with the results by Jaklitsch and Voglmayr. The new subclade contained *T. hypoxylon*, *T. taxi* and *T. rubi* together readily distinguished from other *Trichoderma* species. *Trichoderma hypoxylon* clearly differed from *T. taxi* and *T. rubi*, resulting the sequences similarity of RPB2 and TEF1-α are less than 97%.

*Trichoderma rubi* was found as a new saprobe on stems of *Rubus ulmifolius*, *T. taxi* was reported as a new endophyte of *Taxus mairei*, whereas *T. hypoxylon* was an inhabitant on stroma of *Hypoxylon anthochrous*. In spite of their ecological niches, it showed closed phylogenetic relationship (Fig. 1, Figs S1–S3) and morphological similarity. However, they are a little different in conidiophores and conidia size, and *T. hypoxylon* did not produce pigment on PDA medium, whereas *T. rubi* produced brownish pigment and yellow crystals on PDA medium.

*Trichoderma* species are a rich source of secondary metabolites, probably resulting their environments adaptation and lifestyles. Comparative genome analysis revealed that fungicidal *T. atroviride* and *T. virens* are enriched in secondary metabolism-related genes compared with the biomass-degrading *Trichoderma reesei*. Chemical analysis also showed that these fungicidal species could produce more peptaibols, peptides, polyketides, pyrones, siderophores, terpenoids/steroids than those non-fungicidal *Trichoderma*. These compounds are ecologically and commercially important for their antimicrobial and anti-cancer properties, as well as their ability to induce systemic resistance in plants against microbial invasion. As a new fungicidal species, the chemical diversity of *T. hypoxylon* aroused our attentions. Therefore, the secondary metabolites of this fungus were evaluated. Four major compounds were characterized (Fig. 3). Trichodermandime and aspergillazines are two kind of modified dipeptides. Trichodermandime A has been found from a marine-derived *T. virens*, it is also be obtained from marine-derived fungi *Spicaria elegans* and *Neosartorya pseudafofischeri* and endophytic fungus *Trichoderma spirale*. The yield of trichodermandime A produced by these fungi were 1.58 mg/L, 0.73 mg/L, 0.12 mg/L and 8.98 mg/L individually. In this study, we found that trichodermandime A was one of major compounds in the terrestrial fungus *T. hypoxylon*. It suggested that *Trichoderma* species are important resource for exploration of trichodermandime. Aspergillazines were firstly reported from a soil fungi *Aspergillus unilatialis* (MST-F867), it showed antibacterial and anti-cancer activity. Aspergillazines A been found from two marine-derived fungi *Spicaria elegans* and *T. virens*, in which the productive rate of aspergillazines A was 3.17 mg/L and 0.47 mg/L. This compound has been found co-occurred with trichodermandime A in *T. virens*. Trichothecenes are a well-studied class of sesquiterpene-based mycotoxins. These compounds are a rich source of secondary metabolites, probably resulting their environments adaptation and lifestyles. They are a little different in conidiophores and conidia size, and *T. hypoxylon* did not produce pigment on PDA medium, whereas *T. rubi* produced brownish pigment and yellow crystals on PDA medium.

**Materials and Methods**

**Isolates and specimens.** Samples were collected on 5 May 2014 in Chiang Mai Province, Thailand. *Trichoderma* strains were isolated from the host by single spore isolation as detailed in Chomnunti et al. The holotype is deposited in the Herbarium of Mae Fah Luang University and the isotype in the Herbarium of Mycology, Chinese Academy of Science. Ex-type living cultures are deposited in the Culture Collection of Mae Fah Luang University (MFLUCC, Chiang Rai, Thailand) and the China General Microbiological Culture Collection Center (CGMCC, Beijing, China). Faceoffungi and Index Fungorum numbers are registered as explained in Jayasiri et al. and Index Fungorum.

**Morphological characterization.** Methods and morphology were described basically following counterparts by Jaklitsch & Voglmayr. Colony radius and characteristics were determined on PDA as detailed in Manamgoda et al. and cornmeal dextrose agar (CMD; Difco cornmeal agar + 2% w/v dextrose) and a defined low nutrient agar (SNA) at 20, 25 and 30 °C in darkness for 7 d, then exposed to artificial light to stimulate conidiation formation until 10 d. Microscopic observations and measurements were made from preparations mounted in 50% lactic acid. Photographs were taken with a Nikon DS-FI2 CCD (Nikon, Japan) connected to a Nikon 80i microscope (Nikon, Japan) for anatomical structures. The statistics presented here are based on measurement of 30 mature conidia (± S.E.) and 30 phialides (± S.E.) at 100 × magnification. To assess and describe their structure and morphology of conidiophores were taken from the edge of conidiogenous pustules or fascicles. Conidia were studied from cultures after 10 d of incubation.

**DNA extraction, PCR amplification and sequencing.** Approximately 50 mg of fungal material from each culture was placed in 600 μL of 2% CTAB buffer and ground with a plastic pestle. Genomic DNA was extracted using a modified CTAB extraction protocol. Three primer pairs, ITS5 and ITS4, rRPB2–7ca and rRPB2–7ca, EF983F and EF2218R were separately used to amplify fragments of ITS, RPB2 and TEF1-α. Each PCR was performed on a Votix thermal cycler (Bio-Rad, CA, USA) using easy tag (Tiangen, Beijing, China) in a final volume of 50 μL containing 10 μmol of each primer and 2 μL of DNA (10 ng/L). Reactions were run with positive and negative controls to ensure accuracy and to detect contamination. Automated sequencing was performed by Sino Geno Max Co. (Beijing, China). The sequences used in this study are deposited in GenBank under the accession number provided in Table S1.

**Phylogenetic analysis.** The ITS and TEF1-α data sets used sequences of isolates CGMCC 3.17906, CGMCC 3.17907 and reference sequences were downloaded from GenBank (Table S1). The ITS, TEF1-α and RPB2 data sets were aligned by MAFFT ver.7.03 using the Q-INS-I strategy, individually. The ambiguous areas
of alignment were located and removed using Gblocks 0.91b. Previous phylogenetic analysis of ITS, RPB2 and TEF1-α sequence data from 260 Trichoderma species was conducted separately with *N. berolinensis* and *N. eustromatica* as outgroup. Then, single and combined genes analyses of ITS, TEF1-α and RPB2 sequence data of 60 phylogenetically closed *Trichoderma* species in Brevicompactum, Deliquescent, Hypoceanum, Longibrachiatum, Polysporum, Psychrophila and green spore sections were carried out. *N. berolinensis* and *N. eustromatica* were arranged as outgroup taxa.

Maximum parsimony (MP) analysis was conducted by PAUP 4.0b104 using a heuristic search with tree-bisection-reconnection branch swapping. All characters were treated as unordered and unweighted, gaps were treated as missing data, sequences were auto-increased and Maxtrees was 5,000. Topological confidence of resulted trees was tested by bootstrap proportion with 1,000 replicates, each with 100 replicates of random addition. Bootstrap proportion (BP) higher than 50% from maximum parsimony analysis from PAUP are given.

Maximum-likelihood (ML) analysis was performed in RAxML45 implemented in raxml GUI v.1.3.46. GTR+GAMMA was specified as the model. The analysis was run with a rapid bootstrap analysis using a random start with rapid bootstrap analysis with 1,000 replicates. Bootstrap proportion (BP) higher than 50% from maximum likelihood analysis from RAxML are given.

Bayesian Inference (BI) analysis was performed with MrBayes 3.1.2 using Markov chain Monte Carlo (MCMC) algorithm. Appropriate nucleotide substitution models was determined by MrModeltest 2.349 and the best fit model “GTR + I + G” was selected by Akaikie Information Criterion for the investigated data set. Six chains (one cold and three heated) of 135,000 Markov chain Monte Carlo generations were run, sampling every 100 generation resulting in 1,350 total trees (in two simultaneous analyses). The initial 337 trees (25%) were discarded as burn-in phase of the analyses, and the remaining trees in each analysis were used to calculate posterior probabilities (PP) in the majority rule consensus tree49, posterior probabilities greater than 0.95 are given.

All trees were viewed in TreeView 1.6.640 and revised in Adobe Illustrator CS5.

**Analytical methods and equipment overview.** 1H-NMR spectra were recorded on a Bruker Avance-500 spectrometer using TMS as internal standard, and chemical shifts were recorded as δ values. ESI-MS utilized on an Agilent Accurate-Mass-QTOF LC/MS 6520 instrument. HPLC analysis was performed on a Waters HPLC system (Waters e2695, Waters 2998, Photodiode Array Detector) using an ODS column (C18, 250 × 4.6 mm, YMC Pak, 5 μm) with a flow rate of 1 mL/min.

**Fermentation and isolation.** The fungal strain was cultured on 20 slants of potato dextrose agar at 25 °C for seven days. The fermented PDA substrate was extracted repeatedly with ethyl acetate by exhaustive maceration (4 × 200 mL). The organic solvent was evaporated to dryness under vacuum to afford the crude extract (330 mg). The residue was fractionated by Sephadex LH-20 CC using CH3Cl:Acetone (V/V = 1:1) elution to obtain fifteen fractions 1–15. The fractions 3–7 (200 mg) was separated by semi-preparative RP-HPLC (Waters PrepTM C18 column; 7 μm; 7.8 × 300 mm) with a flow rate of 1 mL/min.

**Trichodermae A (1).** White powder (MeOH) 23,26; 1H NMR (500 MHz, in Acetone-d6) δ 2.72 (dd, J = 19.4, 2.2, H-3a); 2.28 (dd, J = 19.4, 3.9, H-3b); 4.48 (m, H-5); 5.60 (ddd, J = 10.4, 2.0, 2.0, H-6); 5.55 (ddd, J = 10.4, 2.0, 2.0, H-7); 4.24 (m, H-8); 4.15 (dd, J = 7.8, 2.2, H-9); 8.58 (s, H-3′); 7.41 (dd, J = 8.8, H-5); 7.12 (dd, J = 8.8, H-6); 3.96 (s, 7′-OCH3); 3.91 (s, 8′-OCH3); 9.40 (s, CONH); Positive ESIMS: m/z 433.1 [M+H]+.

Aspergillazine A (2). Yellow powder (MeOH) 26,27; 1H NMR (500 MHz, in Acetone-d6) δ 3.14 (d, J = 11.7, H-3a); 2.42 (d, J = 11.7, H-3b); 4.16 (d, J = 4.9, H-5); 5.95 (dd, J = 10.0, 4.9, H-7); 6.07 (dd, J = 10.0, 4.9, H-7); 4.33 (m, H-8); 4.23 (br s, H-9); 6.90 (s, H-3′); 7.09 (dd, J = 8.8, H-5); 6.69 (d, J = 8.8, H-6); 3.90 (s, 7′-OCH3); 3.83 (s, 8′-OCH3); 9.38 (s, CONH); 9.90 (s, -O-NH); Positive ESIMS: m/z 499.1 [M+H]+.

Aspergillazine C (3). Yellow powder (MeOH) 27; 1H NMR (500 MHz, in Acetone-d6) δ 2.96 (d, J = 13.9, H-3a); 2.40 (d, J = 13.9, H-3b); 4.31 (m, H-5); 5.42 (dd, J = 10.0, 1.8, H-6); 5.49 (dd, J = 10.0, 1.9, H-7); 4.16 (dd, J = 7.8, 1.2, H-8); 3.77 (dd, J = 8.0, H-9); 6.85 (s, H-3′); 7.11 (d, J = 8.8, H-5); 6.69 (d, J = 8.8, H-6); 3.90 (s, 7′-OCH3); 3.84 (s, 8′-OCH3); Positive ESIMS: m/z 451.1 [M+H]+.

Harzianum B (4). White powder (MeOH) 28; 1H NMR (500 MHz, in CDCl3) δ 3.84 (d, J = 5.2, H-2); 2.57 (dd, J = 15.5, 7.8, H-3a); 2.03 (ddd, J = 15.5, 5.3, 3.6, H-3b); 5.65 (dd, J = 7.8, 3.6, H-4); 1.92–1.96 (m, H-7a); 1.40–1.43 (m, H-7b); 1.97–1.99 (m, H-8); 5.41 (d, J = 5.8, H-10); 3.63 (d, J = 5.8, H-11); 3.13 (d, J = 4.0, H-13a); 2.83 (d, J = 4.0, H-13b); 0.72 (s, H-14); 0.95 (s, H-15); 1.71 (s, H-16); 6.03 (d, J = 15.4, H-2′); 7.32 (dd, J = 15.4, 10.2, H-3′); 6.61 (dd, J = 10.2, 10.2, H-4′); 6.64 (dd, J = 10.2, 10.2, H-5′); 7.32 (dd, J = 15.4, 10.2, H-6′); 6.07 (d, J = 15.4, H-7′); Positive ESIMS: m/z 401.2 [M+H]+.

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Author Contributions
X.L., W.Y. and K.H. initiated and coordinated the project, and also edited the manuscript. J.S. isolated the fungus and did the morphology and phylogenetic analyses and initially wrote the manuscript. Y.P. extracted genomic DNA and amplified genes. E.L. and W.L. studied the metabolites.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep

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