Community structure of gut fungi during different developmental stages of the Chinese white pine beetle (*Dendroctonus armandi*)

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The Chinese white pine beetle is arguably the most destructive forest insect in the Qinling Mountains in Northern China. Little is known about the structure of the fungal communities associated with *Dendroctonus armandi*, even though this wood-boring insect plays important roles in ecosystem and biological invasion processes that result in huge economic losses in pine forests. The aim of this study was to investigate the fungal community structure present in the galleries and guts of *D. armandi* at different developmental stages using PCR-denaturing gradient gel electrophoresis (DGGE). Analysis of PCR-amplified 18S rRNA gene fragments of fungi from the guts of *D. armandi* revealed fungal communities of low complexity that differed according to the developmental stage. Yeast of the genus *Candida* and the filamentous fungi *Ophiostoma* predominated in *D. armandi* and its galleries. In particular, *Candida* accounted for 56% of the fungal community in the pupal stage. Characterizing the evolution and content of the intestinal microbial community structure in *D. armandi* may facilitate the development of new pest control strategies.

Bark beetles are among the most ecologically and economically important insects in forests worldwide. Their enormous influence in these ecosystems is enabled by microbial alliances that support their exploitation of trees\(^1\). Most bark beetle species engage in symbiotic relationships with fungi\(^2,3\) and rely on these fungi to overcome their limited metabolic abilities\(^4,5\). Bark beetles of the weevil subfamily Scolytinae increase their host-colonizing potential through symbiotic relationships with fungi present in specialized structures called mycangia or on the insect body surface\(^2,7\). Some beetles supplemented their nitrogen-poor phloem diet with fungi (mycophloecophagous) growing in their walls\(^8,9\). View the knowledge of the bark beetle holobiont discovered to date, it is found that beetles rely on microbes to perform basic life functions and to exploit resources and habitats\(^10,11,12\). Thus, associations in bark beetle-fungus systems are of particular research interest. However, on this topic have been mainly focused on a relatively few North American tree-killing beetles. To better understand these symbioses, studies focusing on interactions between beetles and microbes from other regions of the world are needed.

Bark beetles, particularly *Dendroctonus* species, are serious pests in coniferous forests and cause large economic losses\(^13\). The Chinese white pine beetle (*Dendroctonus armandi* Tsai and Li, Scolytidae) kills living *Pinus armandi* and has caused serious damage to *P. armandi* forests in the Qinling and Bashan Mountains in Northern China since 1954\(^14\). Various aspects of the biology and physiology of *D. armandi* have been studied, including its niche within the *P. armandi* ecosystem\(^14\), its life cycle\(^15\), and its symbiotic microbes\(^16,17\). We previously investigated the symbiotic blue stain fungus present in the mycangia of *D. armandi*\(^16,18,19\), but studies of the symbiotic intestinal fungal communities of *D. armandi* have been limited.

Fungal communities play important roles in the life histories of bark beetles\(^7\). The fungi associated with the symbiotic complex may be compared to a ‘root system’ or an ‘external stomach’ of the insect host due to the ability of these fungi to concentrate large amounts of phloem nutrients\(^20\). Intimate associations of ascomycetous and basidiomycetous fungi with phytophagous Coleoptera have been reported including not only mutualistic but also commensals and antagonists relationships\(^21,22\). Experimental evidence suggests that certain yeasts in the alimentary canals of bark beetles might be involved in digestive and detoxification processes and in the production of pheromones essential for beetle chemical communication\(^23,24\). Some gut-associated microflora exhibit spatial and geographic variation\(^24,25\), but their potential variation among development stages and between sexes is poorly
understood. In general, defining community membership is the first step toward understanding the roles of community members in bark beetle biology and in the functioning of the community itself.

We previously investigated the evolution of gut-associated bacterial communities in the ontogeny of D. armandi\(^v\). In the present study, to obtain accurate information on the microbial diversity of D. armandi, the gut-associated fungal communities of D. armandi were investigated in the larval, pupal, and adult stages and their galleries in the host P. armandi using a culture-independent method (i.e., DGGE). We determined which fungal species were consistently associated with the Chinese white pine beetle throughout all developmental stages.

### Results

**Fungal diversity analysis.** The number, density, and composition of fungal DGGE bands were different at each development stage (Fig. 1). Based on the peak density of bands in the DGGE profile, fungal diversity indices were analyzed to estimate the diversity of the microbial communities, i.e., the value of the Shannon-Wiener index ($H'$) was positively related to the diversity of the fungal community. The richness and $H'$ value (2.996725) of the fungi were highest in D. armandi larval guts (S4–S6), with 21 visible bands in the DGGE profiles, and lowest in the pupal stage (S7–S9), with 12 visible bands (Table 1). ANOVA confirmed that there was a significant difference in fungal diversity ($H'$) between the galleries (S1–S3) and all D. armandi development stages ($P < 0.05$), except for the pupal stage ($P > 0.05$). The $H'$ value was significantly different among all development stages (S4–S15) ($P < 0.05$).

**Phylogenetic analyses and dominant taxa.** In total, 19 different bands were successfully sequenced, and a phylogenetic tree was constructed (Fig. 2). According to the identification results of the 19 bands, it was found that most of the species were related to Ascomycota (C1–C10, C12–C18), with 11 species (C1–C10, C17) belonging to Saccharomycetales. Additional taxa were C11 (Zygomyctys, Mortierella) and C19 (Basidiomycota, Cryptococcus).

The peak density of band C9 was highest in lanes S1–S12, whereas band C19 displayed the highest peak density in lanes S13–S15 (Fig. 1). The phylogenetic analysis indicated that C9 exhibited a high degree of similarity (99% bootstrap support) to Candida ernobii and Candida karawaiewii, and C19 exhibited 99% similarity to Cryptococcus carnescens. That is, Candida sp. represented the most dominant species in the galleries, larvae, pupae, and female adults, whereas Cryptococcus carnescens was the most frequent species among the male adults.

Analysis at the genus level identified abundant sequences matching those of Candida, Kuraishia, Myxozyma, Ophiostoma, Mortierella, and Cryptococcus (Fig. 3). The most predominant fungal genus in the galleries and guts of D. armandi was Candida at all development stages. In the pupal stage, the percentage of Candida reached 55.9% (Fig. 3). Yeasts of the genus Kuraishia were present in all samples except for the pupae. The genus Mortierella was present in all samples except for the male adults, whereas Cryptococcus was only present in the male adults (Fig. 3).

**Operational Taxonomic Units (OTUs).** The 97% sequence similarity level, corresponding to OTUs\(_{0.03}\), was used in the subsequent analyses. Based on the distance matrices of the sequences, nine OTU\(_{0.03}\) clusters were observed in the fungal communities of all the tested samples, including the galleries and different developmental stages of D. armandi (Fig. 2). The sequences belonging to clusters I and IV were grouped into the same phylogenetic branch and are closely related to Candida (99% bootstrap support) and Myxozyma (99% bootstrap support), respectively, cluster VIII is related to Ophiostoma (98% bootstrap support) (Fig. 2). In addition, cluster I was the most dominant group among the samples.

The Chao index was used to better characterize the relative fungal species richness in the different stages and sexes of D. armandi. According to the OTU and richness estimation results, the simplest fungal community structure occurred in the pupal gut (Fig. 4), with five OTU\(_{0.03}\) clusters (I, IV, V, VII, and VIII) belonging to Ascomycota and Zygomyctys (Fig. 2). The larval fungal species richness was the highest among all the developmental stages (S4–S6), with seven OTU\(_{0.03}\) clusters (I, II, and IV–VIII) related to Ascomycota and Zygomyctys (Fig. 2). The galleries, female and male samples, including the galleries and different developmental stages of D. armandi (Fig. 2). The sequences belonging to clusters I and IV were grouped into the same phylogenetic branch and are closely related to Candida (99% bootstrap support) and Myxozyma (99% bootstrap support), respectively, cluster VIII is related to Ophiostoma (98% bootstrap support) (Fig. 2). In addition, cluster I was the most dominant group among the samples.

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![Figure 1](https://www.nature.com/scientificreports/)

**Figure 1** | DGGE profiles of nested PCR-amplified 18S rRNA gene fragments of fungi from the galleries and guts of Dendroctonus armandi in different development stages. Lanes S1 to S3 correspond to DGGE profiles of three galleries samples, S4 to S6 correspond to DGGE profiles of three D. armandi larvae samples, S7 to S9 correspond to DGGE profiles of three D. armandi pupae samples, S10 to S12 correspond to DGGE profiles of three female D. armandi adults samples, and S13 to S15 correspond to DGGE profiles of three male D. armandi adults samples. Bands C1 to C19 represent 18S rRNA gene regions of different fungi.
Table 1 | Richness (S), evenness (E_H), and Shannon-Wiener index (H') of intestinal symbiotic fungi identified in galleries and guts of *Dendroctonus armandi* at different developmental stages

| Lane  | Sample     | S       | E_H     | H'      |
|-------|------------|---------|---------|---------|
| S1–S3 | Galleries  | 12      | 0.966 ± 0.012 | 2.401 ± 0.031 |
| S4–S6 | Larvae     | 21      | 0.984 ± 0.002 | 2.997 ± 0.005 |
| S7–S9 | Pupae      | 12      | 0.976 ± 0.001 | 2.419 ± 0.003 |
| S10–S12 | Female adults | 18   | 0.981 ± 0.002 | 2.833 ± 0.005 |
| S13–S15 | Male adults | 16    | 0.944 ± 0.003 | 2.615 ± 0.007 |

Note: Data are means ± standard error (SE), n = 3.

Figure 2 | Neighbor-joining phylogenetic tree of the fungal community in the guts of *Dendroctonus armandi* and galleries from their host based on 18S rRNA gene fragments with the model Kimura 2-parameter + G. I–IX represent 9 different OTU_{0.03} clusters obtained using MOTHUR.
adult libraries each included six OTU$_{0.03}$ clusters according to the Chao index (Fig. 2). In the guts of female adults, six clusters (I, III, IV, and VI–VIII) were related to Ascomycota and Zygomyxota; in male adults, these clusters were I, III–V and VIII–IX, which belong to Ascomycota and Basidiomycota.

**Discussion**

Bark beetles are associated with an array of filamentous fungi and yeasts. In our study, gut-associated filamentous fungi and yeasts were surveyed in *D. armandi* beetles and galleries. The present study provides the first insight into the gut fungal diversity of *D. armandi* at different developmental stages and among the two sexes. Overall, the results of the fungal diversity indices constructed based on the peak density of bands in the DGGE profiles were consistent with the richness estimation according to OTUs. The predominant species of the fungal community in *D. armandi* formed a group of low complexity, the structure of which differed among the developmental stages. With the exception of fungus-feeding beetles, this low level of fungal complexity is typical of the bark beetle gut discovered to date. The diversity of gut-associated fungi in conifer-associated bark beetles is also relatively low, in which only eight fungal species from three genera were identified on the body surfaces of *Dendroctonus ponderosae* larvae, pupae, and adults. Incorporating both culture-based and molecular methods, only 14 yeast species in five genera were identified on the surfaces of *D. ponderosae* Hopkins and *Ips pini* (Say). Similarly to previously reported associated bacteria, some fungi were conserved in the galleries and guts of all *D. armandi* stages studied, although not always in the same abundance, including taxa of clusters I (*Candida*) and V (*Myxozyma*) related to yeasts and VIII (*Ophiostoma*) related to the blue stain fungus (Fig. 2). This result...
indicates that these groups are consistently associated with *D. armandi* or their host trees.

Yeasts are commonly associated with bark beetles and might be an important nutritional source for the insect host. Furthermore, yeasts might contribute to successful brood development by limiting the numbers of beetles in individual trees, as some yeast can convert the bark beetle’s aggregation pheromone trans-verbenol into the anti-aggregation pheromone verbenone. Yeasts that are consistently present in the *D. armandi* gut at different developmental stages might influence the development of the insect host.

Ascomycetes of the genus *Ophiostoma*, also known as ‘blue stain fungi’, are typically associated with conifers. These fungi produce asexual and sexual spores in slimy masses that attach to insect bodies and are dispersed to new hosts that represent fresh nutrient sources. When bark beetles invade conifers, the fungus taps into the sapwood and transports it to the phloem and bark where the beetle larvae feed, increasing the nitrogen content by up to 40%; this is critical for bark beetle development and survival. *Ophiostoma* taxa are also the most important filamentous fungal associates of the mountain pine beetle, according to the results of a survey of its body surface, including the mycangia; however, the effects of these fungi on beetles as gut associates are mostly unknown. To further understand the *Ophiostoma-Dendroctonus* relationship, further studies are needed.

The results of the DGGE analyses indicated that Ascomycota species are prevalent in both *D. armandi* beetles and their galleries. In the combined diversity indices, Ascomycota belonging to the genus *Candida* was the most abundant taxon. *Candida* is ubiquitous in the environment and responsible for the majority of all fungal infections worldwide. *Candida* species have also been isolated from the guts of other *Dendroctonus* spp. Suh et al. reported several examples of highly specific associations between certain beetles and yeasts. Our results support this result because certain yeasts were present during the entire life cycle of their beetle hosts. In addition, *Candida ernobii* is able to produce lipase, which catalyzes the hydrolysis of long-chain triacylglycerols and plays an important physiological role in several metabolic processes (fat digestion in the gastro-intestinal tract, lipolysis in the adipose tissue, lipolysis of lipoproteins) in the conversion of oils and fats into free fatty acids and partial acylglycerols. *Candida ernobii* was the most abundant gut associate and might play similar roles in *D. armandi*. Further experiments are required to verify the roles of yeast, particularly *Candida* spp., in the guts of *D. armandi*.

Among the tested developmental stages, the pupal stage was characterized by the simplest fungal structure according to both diversity indices and the Chao index. Low microbial diversity in the pupal gut is common in bark beetles. Energy is required to achieve metamorphosis through the activation of several metabolic processes and morphological changes, even though pupae do not feed. Therefore, the decrease in fungal diversity and increase in *Candida* spp. in pupae observed in this study could be associated with the absence of feeding activity and the necessity for pupae to obtain energy more efficiently by adjusting their microbial community structure.

Attempts have been made to use fungi as biocontrol agents against bark beetle pests in forestry biological control practices. Wegensteiner has presented an overview of the entomopathogenic fungal encountered in European bark beetle galleries. Although few studies have examined the utilization of gut fungi to control bark beetles, gut fungi are candidates for future pest biocontrol strategies. However, detailed knowledge about the dynamic variation, colonization, and modes of transmission of intestinal fungi is required before such a strategy could be successfully implemented.

The information about fungal community structure collected in this study will provide the basis for subsequent studies on the roles of these fungal associates in bark beetle development, ecology, and management. Future studies confirming and quantifying the ability of fungi to degrade tree defenses are required to understand the influences of these fungi on bark beetle biology.

### Methods

#### Study sites.

The material collection site was the Huodaping Experimental Forest Station of the Northwest Agricultural & Forestry University. The station is located on the southern slope of the middle Qinling Mountains (33° 18′–33° 28′N, 108° 21′–108° 39′E), Shaanxi, China. *Pinus armandi*, *Picea wilsonii*, and *Pinus tabulaeformis* are the common tree species distributed throughout most parts of the Qinling Mountains.

#### Insect collection and dissection.

Larvae, pupae, and female and male adults of *D. armandi* and 0.5 × 2-cm sections of galleries were removed from recently attacked *P. armandi* phloem and placed on ice before transport to the laboratory. All *D. armandi* samples were manually obtained directly from galleries of infested pine trees using centrifugal and were then transferred to a separates the microvials containing sterile moist paper. To investigate the influence of sex on gut-associated fungi, female and male adults were separated according to their reproductive organs. For DGGE analysis, 120 beetle samples in each developmental stage and gallery samples were obtained from attacked *P. armandi*. 40 beetle samples in each stage were pooled to be one sample respectively. A total of five samples representing galleries and different beetle developmental stages were obtained. The experiment was repeated twice.

The insect samples were rinsed with sterile water, surface sterilized with 70% ethanol for 3 min, and again rinsed twice with sterile water. After placing in 10 mM sterilized phosphate-buffered saline (138 mM NaCl and 2.7 mM KCl pH 7.4), the samples were dissected under a stereomicroscope using insect pins to obtain mid-guts and hindguts. Forty guts from each life stage were transferred to 1.5-mL micro-centrifuge tubes and then homogenized several times with a plastic pestle in liquid nitrogen, followed by vortexing with 500 µL of Tris-EDTA [10 mM Tris-HCl (pH 8.0), 1 mM EDTA] for 3 min at the speed of 2500 rpm. The homogenate was centrifuged at 10,000 rpm for 30 s to separate the gut wall tissues and undigest food. The supernatant (containing fungi) was transferred to new tubes for DNA extraction. All procedures were completed in a sterile environment (Biological Air Clean Bench, Suzhou Antai Airtech, Jiangsu, China).

#### Fungal DNA extraction.

Fungal DNA was extracted using the E.Z.N.A. Fungal DNA Kit (Omega, Biotech, Doraville, GA, USA) according to the manufacturer’s directions and stored at −20°C until use.

#### Nested PCR.

Nested PCR was used to increase the resolution of DGGE. The highly variable region of the fungal 18S rRNA gene was amplified using the primer pairs NS1/EF3 and FF390/FR1-GC. All PCR amplifications were performed in an S1000™ Thermal cycler (Bio-Rad, Hercules, CA, USA) in a final volume of 50 µL containing 25 µL of 2X Taq Master Mix (CoWin Biotech, Beijing, China), 0.8 µL of each primer (10 µM, Invitrogen Trading, Shanghai, China), 1 µL of template, and 22.4 µL of RNase-free water. The fungal DNA was diluted to 30 ng µL−1 and used as the template for first-round PCR, which was performed using the following program: 94°C for 3 min, followed by 30 cycles of 94°C for 50 s, 56°C for 45 s, and 72°C for 50 s, and 72°C for a final extension at 72°C for 5 min. The product of the first-round PCR was diluted 1/200 with ddH2O and further deployed as the template in the second PCR reaction, which was performed using the following procedure: 94°C for 3 min, followed by 35 cycles of 94°C for 30 s, 58°C for 30 s, and 72°C for 30 s and a final step at 72°C for 5 min.

The PCR products of DNA extraction, primer specificity were analyzed by PCR electrophoresis and ethidium bromide staining in the presence of the DL2000 DNA marker (Takara Biotechnology, Dalian, China). The obtained PCR products were stored at −20°C until DGGE analysis.

#### Denaturing gradient gel electrophoresis (DGGE).

The DCode™ Universal Mutation Detection System (Bio-Rad, Hercules, CA, USA) was used for the DGGE analysis. A 35-µL aliquot of fungal nested-PCR product per sample was loaded onto an 8% (w/v) poly-acrylamide (37.5: 1 acrylamide/bio-acrylamide) gel containing a linear denaturing gradient of 40% to 60%, where 100% denaturing acrylamide contained 7 M urea and 40% formamide. The gel was initially electrophoresed at 120 V for 10 min, followed by electrophoresis at 70 V for an additional 1 h at 58°C in 1× TAE buffer [40 mM Tris-acetate (pH 7.4), 20 mM sodium acetate, 1 mM disodium EDTA]. After staining with ethidium bromide solution for 10 minutes, the gel was destained in deionized water for 10 minutes and then photographed under UV light using the Gel Doc™ XR System (Bio-Rad, CA, USA).

#### DGGE band identification.

The dominant DGGE bands were excised from the polyacrylamide gel, and the DNA was eluted using an E.Z.N.A. Mag-Bind Poly-Gel DNA Extraction Kit (Omega, USA). The eluted DNA (1 µl) was re-amplified with primer pair FF390 and FR1 without a GC clamp at the 5′ end. The PCR products were purified using a Gel Extraction Kit (Baitaile Biological Technology, Xi’an, China) and inserted into the pMD18-T vector (pMD18-T Cloning Kit, Takara Biotechnology, Dalian, China). The recombinant plasmids were transformed into Escherichia coli (strain DH5α), and positive colonies were identified by blue-white screening. Five positive colonies were randomly selected from each transformation to further confirm the presence of correct inserts via PCR with the primer pair M13-47 and 51-21.
M13–48. The confirmed clones were sequenced (Jinsirui Biotechnology, Nanjing, China).

To classify the intestinal fungi, the obtained sequences were matched to those in the RDP II database and searched in the NR database to select and download reliable and highly similar sequences (≥97% bootstrap support). The sequences obtained in this study have been submitted to the NCBI database under the accession numbers KF928180–KF928181. The phylogenetic relationships of the intestinal fungi were analyzed by molecular phylogeny. The sequences were aligned using MUSCLE, available in the software MEGA 5.2, and the best model was computed. The phylogenetic trees were constructed using the maximum likelihood and neighbor-joining methods. To calculate the support for each clade, a bootstrap analysis was performed with 1000 replications.

**DGGE band profile analysis.** The DGGE band profiles were analyzed using Quantity One software (Bio-Rad, USA) by the following procedure. First, auto frame lanes were selected, and the rolling disk size was adjusted to 5 to minimize the influence of background. Second, the bands were detected, and the parameters were adjusted to acquire the most reliable band pattern; the Gauss model was applied to all lanes. Third, the lane with the most bands was selected for auto-match, and the tolerance was set at 4.00%; the other lanes were matched manually. Finally, the peak density of all lanes was reported for further analysis.

Each band was digitized via auto-detection of the peak density. Based on the transferred data, diversity indices were calculated to investigate the dominant fungal communities and to determine their changes in the galleries, larvae, pupae, and adults. The DGGE band profile analysis was performed with 1000 replications.

2. Paine, T. D., Raffa, K. F. & Harrington, T. C. Interactions among scolytid bark beetles. *Science*. 276, 63–66 (1999).

3. Hu, X., Wang, C., Chen, H. & Ma, J. Differences in the structure of the gut bacteria communities in development stages of the Chinese white pine beetle (*Dendroctonus armandi*). *Int. J. Mol. Sci.* 14, 21006–21020 (2013).

4. Chen, H., Tang, M., Zhu, C. J. & Hu, J. Y. The enzymes in the secretions of *Dendroctonus armandi* (Scolytidae) and their symbiotic fungi of Leptographium guilliermondii. *Science* 289, 223–234 (2010).

5. Gibson, C. M. & Hunter, M. S. Extraordinarily widespread and fantastically complex: comparative biology of endosymbiotic bacterial and fungal mutualists of insects. *Ecol. Lett.* 13, 223–234 (2010).

6. Six, D. L. The bark beetle holobiont: why microbes matter. *J. Chem. Ecol.* 39, 989–1002 (2013).

7. Six, D. L. & Paine, T. D. Integrative management of bark beetles. *Annu. Rev. Entomol.* 57, 197–206 (2012).

8. Jansson, J. K., Strome, J. O., Singer, M. S. & Abbot, P. Phytophagous insect-microbe mutualisms and adaptive evolutionary diversification. *Evolution.* 62, 997–1012 (2008).

9. Gibson, C. M. & Hunter, M. S. Extraordinarily widespread and fantastically complex: comparative biology of endosymbiotic bacterial and fungal mutualists of insects. *Ecol. Lett.* 13, 223–234 (2010).

10. Six, D. L. Bark beetle-fungus symbioses. *Insect symbiosis.* 1, 97–114 (2003).

11. Vega, F. E. & Dowd, P. F. The role of yeasts in insect endosymbiosis. *Insect-Fungal Associations: Ecology and Evolution.* Vega, F. E. & Blackwell, M. (eds.) 211–243 (Oxford University Press, New York, 2005).

12. Brackwell, R. R. & Six, D. L. Broadscale specificity in a bark beetle–fungus symbiosis: a spatio-temporal analysis of the mycangial fungi of the western pine beetle. *Microb. Ecol.* 68, 859–870 (2014).

13. Coulson, R. N. & Stark, R. W. Integrative management of bark beetles. *Bark beetles in North American forests.* Mitton, J. B. & Sturgeon, K. B. (eds.) 315–349 (University Press of Texas, Austin, 1982).

14. Chen, H. & Tang, M. Spatial and temporal dynamics of bark beetles in Chinese white pine in Qinling Mountains of Shaanxi Province, China. *Environ. Entomol.* 36, 1124–1130 (2007).

15. Yin, H., Huang, F. & Li, Z. Economic insect fauna of China. Fasc. 29. *Coleoptera Scolytidae.* Yin, H., Huang, F. & Li, Z. (eds.) 26–35 (Science Press, Beijing, China, 1984).

16. Ming, T. & Hui, C. Effect of symbiotic fungi of *Dendroctonus armandi* on host trees. *Science*. 35, 63–66 (1999).

17. Hu, X., Wang, C., Chen, H. & Ma, J. Differences in the structure of the gut bacteria communities in development stages of the Chinese white pine beetle (*Dendroctonus armandi*). *Int. J. Mol. Sci.* 14, 21006–21020 (2013).

18. Chen, H., Tang, M., Zhu, C. J. & Hu, J. Y. The enzymes in the secretions of *Dendroctonus armandi* (Scolytidae) and their symbiotic fungi of *Leptographium guilliermondii*. *Science* 289, 223–234 (2010).

19. Nardon, P. & Grenier, A. M. Endocytobiosis in Coleoptera: biological, biochemical, and genetic aspects. *Insect endocytobiosis: morphology, physiology, genetics, evolution.* Schenewerk, W. & Gassner, G. (eds.) 175–216 (CRC Press, Boca Raton, Florida, 1989).

20. Rivera, F. N. et al. Gut-associated yeast in bark beetles of the genus *Dendroctonus Erichson* (*Coleoptera: Curculionidae: Scolytinae*). *Biol. J. Linnean Soc.* 98, 325–342 (2009).

21. Borden, J. H. Aggregation phenonemes. *Bark Beetles in North American Conifers: a system for the study of evolutionary biology.* Mitton, J. B. & Sturgeon, K. B. (eds.) 74–139 (University of Texas Press, Texas, Austin, 1982).

22. Funk, D. J. & Bernays, E. A. Geographic variation in host specificity reveals host range evolution in *Uroleucon ambrosiae* aphids. *Ecology.* 82, 726–739 (2001).

23. Tsuchida, T., Koga, R., Shibao, H., Matsumoto, T. & Fukatsu, T. Diversity and geographic distribution of primary endosymbiotic bacteria in natural populations of the pea aphid, *Acyrthosiphon pisum*. *Mol. Ecol.* 11, 2123–2135 (2002).

24. Broderick, N. A., Raffa, K. F., Goodman, R. M. & Handelmsan, J. Census of the bacterial community of the gypsy moth larval midgut by using culturing and culture-independent methods. *Appl. Environ. Microbiol.* 70, 293–300 (2004).

25. Suh, S. O., Marshall, C. J., Mchugh, J. V. & Blackwell, M. Wood ingestion by passalid beetles in the presence of xylase-fermenting gut yeasts. *Mol. Ecol.* 12, 3137–3147 (2003).

26. Lee, S., Kim, J. J. & Breuil, C. Determining fungal diversity on *Dendroctonus ponderosae* and *Ipis* pinus affecting lodgepole pine using cultural and molecular methods. *Fungal Divers.* 19, 79–94 (2005).

27. Lee, S., Kim, J. J. & Breuil, C. Diversity of fungi associated with mountain pine beetle, *Dendroctonus ponderosae*, and infested lodgepole pines in British Columbia. *Fungal Divers.* 22, 91–105 (2006).

28. Whitney, H. Relationships between bark beetles and symbiotic organisms. *Bark beetles in North American conifers: a system for the study of evolutionary biology.* Mitton, J. B. & Sturgeon, K. B. (eds.) 183–211 (University of Texas Press, Texas, Austin, 1982).

29. Hunt, D. W. A. & Borden, J. H. Conversion of verbenols to verbenone by yeasts isolated from *Dendroctonus ponderosae* (*Coleoptera: Scolytidae*). *J. Chem. Ecol.* 16, 1385–1397 (1990).

30. Masuya, H., Yamaoka, Y., Kaneko, S. & Yamaura, Y. Ophiostomatoid fungi isolated from Japanese red pine and their relationships with bark beetles. *Mycoscience.* 50, 212–223 (2009).

31. Harrington, T. C. Diseases of conifers caused by species of *Ophiostoma* and *Leptographium*. *Ceratocystis and Ophiostoma: taxonomy, ecology, and pathogenicity.* Wingfield, M. J., Seifert, K. A. & Webber, J. F. (eds.) 161–172 (APS Press, Saint Paul, 1993).

32. Six, D. L. & Paine, T. D. Effects of mycangial fungi and host tree species on progeny survival and emergence of *Dendroctonus ponderosae* (*Coleoptera: Scolytidae*). *Ecol. Entomol.* 27, 1393–1401 (1998).

33. Lee, S., Kim, J. J. & Breuil, C. Diversity of fungi associated with mountain pine beetle, *Dendroctonus ponderosae* and infested lodgepole pines in British Columbia. *Fungal Divers.* 22, 91–105 (2006).

34. Vehrin, K., Haase, G., Kurtzman, C., Jo, J. & Stender, H. Differentiation of *Candida albicans* and *Candida dubliniensis* by fluorescent in situ hybridization with peptide nucleic acid probes. *J. Clin. Microbiol.* 39, 4138–4141 (2001).

35. Suh, S. O. & Blackwell, M. Three new beetle-associated yeast species in the *Pichia* giihermondii clade. *FEMS. Yeast. Res.* 5, 87–95 (2004).
38. Aloulou, A. et al. Purification and biochemical characterization of the LIP2 lipase from Yarrowia lipolytica. Biochim. Biophys. Acta. Mol. Cell. Biol. Lipids. 1771, 228–237 (2007).
39. Morales-Jiménez, J., Zañiga, G., Ramirez-Saad, H. C. & Hernández-Rodríguez, C. Gut-associated bacteria throughout the life cycle of the bark beetle Dendroctonus rhizophagus Thomas and Bright (Curculionidae: Scolytinae) and their cellulolytic activities. Microb. Ecol. 64, 268–278 (2012).
40. Popa, V., Déziel, E., Lavallée, R., Bauce, E. & Guertin, C. The complex symbiotic relationships of bark beetles with microorganisms: a potential practical approach for biological control in forestry. Pest, Manag. Sci. 68, 963–975 (2012).
41. Wegensteiner, R. & Smits, P. Laboratory evaluation of Beauveria bassiana (Bals.) Vuill. and Beauveria brongniartii (Sacc.) Petch against the four eyed spruce bark beetle, Polygraphus poligraphus (L.) (Coleoptera, Scolytidae); Bulletin OILB/SROP 23, 161–166 (2000).
42. Delalibera, I., Handelsman Jr, J. & Raffa, K. F. Contrasts in cellulolytic activities of gut microorganisms between the wood borer, Saperda vestita (Coleoptera: Cerambycidae), and the bark beetle, Ips pini and Dendroctonus frontalis (Coleoptera: Curculionidae). Environ. Entomol. 34, 541–547 (2005).
43. Pittl, E., Innererbner, G., Wanek, W. & Insam, H. Microbial communities of arboreal and ground soils in the Esquinas rainforest, Costa Rica. Plant. Soil. 329, 65–74 (2010).
44. Muyzer, G., De Waal, E. C. & Uitterlinden, A. G. Profiling of complex microbial populations by denaturing gradient gel electrophoresis analysis of polymerase chain reaction-amplified genes coding for 16S rRNA. Appl. Environ. Microbiol. 59, 695–700 (1993).
45. Cole, J. R. et al. The Ribosomal Database Project: improved alignments and new tools for rRNA analysis. Nucleic. Acids. Res. 37, 141–145 (2009).
46. Wheeler, D. L. et al. Database resources of the national center for biotechnology information. Nucleic. Acids. Res. 35, 5–12 (2007).
47. Edgar, R. C. MUSCLE: multiple sequence alignment with high accuracy and high throughput. Nucleic. Acids. Res. 32, 1792–1797 (2004).
48. Tamura, K. et al. MEGAS: molecular evolutionary genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods. Mol. Biol. Evol. 28, 2731–2739 (2011).
49. Galand, P. E., Fritze, H. & Yrjälä, K. Microsite-dependent changes in arbuscular mycorrhizal fungal biodiversity in soil through a land-use gradient using a pyrosequencing approach. Environ. Microbiol. 12, 2165–2179 (2010).
50. Lumini, E., Orgiazzi, A., Burriello, R., Bonfante, P. & Bianciotto, V. Disclosing arbuscular mycorrhizal fungal biodiversity in soil through a land-use gradient using a pyrosequencing approach. Environ. Microbiol. 12, 2165–2179 (2010).
51. Chao, A. & Shen, T. J. Nonparametric estimation of Shannon’s index of diversity when there are unseen species in sample. Ecol. Ecol. Stat. 10, 429–443 (2003).
52. Gotelli, N. J. & Colwell, R. K. Quantifying biodiversity: procedures and pitfalls in the measurement and comparison of species richness. Ecol. Lett. 4, 379–391 (2001).
53. Colwell, R. K., Mao, C. X. & Chang, J. Interpolating, extrapolating, and comparing incidence-based species accumulation curves. Ecology 85, 2717–2727 (2004).

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Author contributions
X.H. and H.C. conceived and designed the experiments. X.H. and M.L. performed the experiments and analysed the data. H.C. provided helps in the experiments. X.H., M.L. and H.C. co-wrote the manuscript. All authors discussed the results and commented on the manuscript.

Additional information
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