Multigene phylogeny, phylogenetic network, and morphological characterizations reveal four new arthropod-associated *Simplicillium* species and their evolitional relationship

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*Simplicillium* species are widely distributed and commonly found on various substrates. A minority of species are associated with arthropods. A spider-associated species *Simplicillium araneae*, and three insect-associated species, *Simplicillium coleopterorum*, *Simplicillium guizhouense*, and *Simplicillium larvatum*, are proposed as novel species based on a multi-locus phylogenetic analysis and morphological characteristics. These *Simplicillium* species completely fit the nutritional model of Hypocreales fungi and could be used as a model to study their evolutionary relationship. A phylogenetic network analysis based on ITS sequences suggests that a host jump was common among *Simplicillium* species, and *S. araneae* may have originally come from an insect host and then jumped to a spider host. However, the evolutionary relationship of *S. coleopterorum*, *S. guizhouense*, and *S. larvatum* was not clear in the phylogenetic network and more sequencing information should be added to the network. In addition, strain CBS 101267 was identified as *Simplicillium subtropicum*.

KEYWORDS
spider, insect, multigene phylogeny, morphological characterization, phylogenetic relationship

Introduction

The genus *Simplicillium* branched off from the genus *Verticillium* section *Prostrata*, and it consists of four species: *S. lanosoniveum* (J.F.H. Beyma) Zare and W. Gams, *S. obelavatum* (W. Gams) Zare and W. Gams, *S. lamellicola* (F.E.V. Sm.) Zare and W. Gams, and *S. wallacei* H.C. Evans (Zare and Gams, 2001). Zare and Gams (2001) summarized that solitary phialides, conidia adhering in globose, slimy heads or imbricate chains, and
crystals commonly present in agar were the typical characteristics of *Simplicillium*. After that, numerous species were added to the genus (Liu and Cai, 2012; Nonaka et al., 2013; Gams, 2017; Zhang et al., 2017; Croux et al., 2018; Gomes et al., 2018; Chen et al., 2019, 2021; Wei et al., 2019; Kondo et al., 2020; Wang et al., 2020; Leplat et al., 2021). However, based mainly on rDNA sequence analyses, several *Simplicillium* species (*S. wallacei* A.A.M. Gomes and O.L. Pereira, *S. chinesis* F. Liu and L. Cai, and *S. filiforme* R.M.F. Silva, R.J.V. Oliveira, Souza-Motta, J.L. Bezerra and G.A. Silva) were transferred to the genera *Lecanicillium* W. Gams and Zare and *Leptobacillium* Zare and W. Gams (Zare and Gams, 2008; Okane et al., 2020; Chen et al., 2021). As a result, the genus *Simplicillium* currently consists of 23 species.

Chen et al. (2021) noted that *Simplicillium* species inhabit diverse substrates and could be used as a model of Hypocreales fungi to study their evolutionary relationship. However, the phylogenetic tree assumes that biological groups evolve in the form of tree divergence and cannot accurately present the whole process of actual evolution, including hybridization, horizontal gene transfer, and gene recombination within the population (Cheng and Huang, 2008). The neighbor-net network (split network), a kind of distance-based phylogenetic network, can be used to present conflicting and ambiguous signals in datasets and detect subtle differences (Bryant and Moulton, 2004; Huson and Bryant, 2006). It can provide a way to present parallel events that are covered up and cannot be displayed by a phylogenetic tree, as well as an uncertain evolutionary phylogenetic relationship. The method has been applied in the phylogenetic analysis of animals, plants, and microorganisms (Bandelt and Dress, 1992; Morrison, 2005; Huson and Bryant, 2006; Morozov et al., 2013; Khonsanit et al., 2020).

During a survey of entomopathogenic fungi from Southwest China, some insect- and spider-associated specimens were found and some new *Simplicillium* strains were isolated and purified. The goals of this research were as follows: (1) identify the new strains based on ITS sequence, (2) characterize the new species of the genus *Simplicillium* based on a multi-locus phylogenetic analysis and their morphological and ecological characteristics, and (3) detect the evolutional relationship of the new species by the neighbor-net network based on ITS sequence of *Simplicillium* species.

### Materials and methods

#### Specimen collection and identification

Five infected insect and spider specimens (DY1005, DY1025, DY10173, DY10181, and SD0538) were collected from Duyun City (26°21’24.71" N, 107°22’48.22" E) and Sandu County (25°57’22.21" N, 107°57’54.69" E), Guizhou Province, on 1 October and 1 May 2019. The surface of each insect body was rinsed with sterile water, followed by surface sterilization with 75% ethanol for 3–5 s and rinsing 3 times with sterilized water. After drying on sterilized filter paper, the synnemata, mycelium, or a part of the sclerotia was removed from the specimen, inoculated on potato dextrose agar (PDA), and improved potato dextrose agar (PDA, 1% w/v peptone) plates (Chen et al., 2019). Fungal colonies emerging from the specimens were isolated and cultured at 25°C for 14 days under 12 h light/12 h dark conditions following protocols described by Zou et al. (2010). The specimens and isolated strains were deposited at the Institute of Fungus Resources, Guizhou University (formally Herbarium of Guizhou Agricultural College; code, GZAC), Guiyang City, Guizhou, China.

Macroscopic characterization was determined from PDA cultures incubated at 25°C for 14 days, and the growth rate of the colony, the presence of octahedral crystals, and the colony colors (surface and reverse) were observed. To investigate the microscopic characteristics, a small number of mycelia were mounted in lactophenol cotton blue or 20% lactate acid solution and observed with an optical microscope (OM, DM4 B, Leica, Germany).

#### DNA extraction, polymerase chain reaction amplification, and nucleotide sequencing

DNA extraction was carried out using a fungal genomic DNA extraction kit (DP2033, BioTeke Corporation) according to Liang et al. (2011). The extracted DNA was stored at −20°C. Amplification of the internal transcribed spacer (ITS) region, large subunit ribosomal RNA (LSU) gene, small subunit ribosomal RNA (SSU), RNA polymerase II largest subunit (RPB1), and translation elongation factor 1 alpha (TEF) was carried out by PCR as described by White et al. (1990), Rakotonirainy et al. (1994), and Castlebury et al. (2004). Primer sequence information is shown in Supplementary Table S1. PCR products were purified and sequenced at Sangon Biotech (Shanghai) Co. The resulting sequences were submitted to GenBank (Table 1).

#### Sequence alignment and phylogenetic and network analyses

Lasergene software (version 6.0, DNASTAR) was used to edit DNA sequences in this study. The SSU, ITS, LSU, RPB1, and TEF sequences were downloaded from GenBank, based on Nonaka et al. (2013), Zhang et al. (2017), Croux et al. (2018, 2021), Gomes et al. (2018), Chen et al. (2019, 2021), Wei et al. (2019), Kondo et al. (2020), Leplat et al. (2021), and others selected based on BLAST searches in GenBank (Table 1). ITS sequence was applied
| Species                      | Strain No. | GenBank accession No. |
|------------------------------|------------|-----------------------|
| **Gamsszerea wallacei**      | CBS 101237 | NR_11267 NG_042398 NG_062646 EF469102 EF469073 |
| **Simplicillium album**      | CGMCC 3.19635 | NR_172844 NG_075278 MK336068 |
| **Simplicillium aogashimaense** | JCM 18167 | AB604002 LC496874 LC496889 LC496904 |
| **Simplicillium aogashimaense** | JCM 18168 | AB604004 LC496875 LC496890 |
| **Simplicillium araneae**    | DY101811   | OMT743774 OMT743792 OMT743793 OMT818465 |
| **Simplicillium araneae**    | DY101812   | OMT743840 OMT743846 OMT743845 OMT818466 |
| **Simplicillium calcicola**  | LC5371     | KU746705 KU746751 KX855251 |
| **Simplicillium calcicola**  | LC5586     | KU746706 KU746752 KX855252 |
| **Simplicillium cicadellidae** | JCM 181101 | MN006243 MN022271 MN022263 |
| **Simplicillium cicadellidae** | JCM 181102 | MN006244 MN022272 MN022264 |
| **Simplicillium coccinellidae** | DY101791 | MT453861 MT453862 MT453863 MT471341 |
| **Simplicillium coccinellidae** | DY101792 | MT453864 MT457410 MT471342 |
| **Simplicillium coleopterorum** | SD05381 | OMT743920 OMT743925 OMT743935 OMT818467 |
| **Simplicillium coleopterorum** | SD05382 | OMT744109 OMT744170 OMT744176 OMT818468 |
| **Simplicillium cylindrosporum** | JCM 18169 | AB603989 LC496876 LC496891 LC496906 |
| **Simplicillium cylindrosporum** | JCM 18170 | AB603994 LC496877 LC496892 LC496907 |
| **Simplicillium cylindrosporum** | JCM 18171 | AB603997 |
| **Simplicillium cylindrosporum** | JCM 18172 | AB603998 |
| **Simplicillium cylindrosporum** | JCM 18173 | AB603999 |
| **Simplicillium cylindrosporum** | JCM 18174 | AB604005 |
| **Simplicillium cylindrosporum** | JCM 18175 | AB604006 |
| **Simplicillium formicidae**  | MFLUCC 18–1379 | MK766511 MK766512 MK765046 MK882623 MK926451 |
| **Simplicillium formicidae**  | DL10041    | MN006241 MN022269 MN022270 |
| **Simplicillium guizhouense** | DY10051    | OMT743225 OMT743226 OMT743242 OMT818453 |
| **Simplicillium guizhouense** | DY10052    | OMT743241 OMT743252 OMT743253 OMT818454 |
| **Simplicillium humicola**    | CGMCC 3.19573 | NR_172845 NG_075279 MK336071 |
| **Simplicillium hymenopterorum** | DY101691 | MT453848 MT453850 MT453849 MT471344 MT471337 |
| **Simplicillium hymenopterorum** | DY101692 | MT453851 MT453853 MT453852 MT471338 |
| **Simplicillium lamellicola**  | CBE 116.25  | AJ292393 AF339552 AF339601 DQ522404 DQ522356 |
| **Simplicillium lamellicola**  | KY00006    | AB378533 AF339552 AF339601 DQ522404 DQ522356 |
| **Simplicillium lamellicola**  | UAMH 2055  | AF108471 AF339552 AF339601 DQ522404 DQ522356 |
| **Simplicillium lamellicola**  | UAMH 4785  | AF108480 AF339552 AF339601 DQ522404 DQ522356 |

(Continued)
to identify the new strains in the genus *Simplicillium* (analysis 1). ITS sequences and other loci were aligned and edited by MAFFT v7.037b ([Katoh and Standley, 2013](#)). Combined sequences of SSU, ITS, LSU, RPB1, and TEF were applied to establish the four novel species (analysis 2) and obtained using SequenceMatrix v.1.7.8 ([Vaidya et al., 2011](#)). The model was selected for Bayesian analysis by ModelFinder ([Kalyaanamoorthy et al., 2017](#)) in PhyloSuite software ([Zhang et al., 2020a](#)). ITS sequences and the combined loci were analyzed using Bayesian inference (BI) and maximum likelihood (ML) methods. For BI, a Markov chain Monte Carlo (MCMC)
algorithm was used to generate phylogenetic trees with Bayesian probabilities using MrBayes v.3.2 (Ronquist et al., 2012) for the combined sequence datasets. The Bayesian analysis resulted in 20,001 trees after 10,000,000 generations. The first 4,000 trees, representing the burn-in phase of the analysis, were discarded, while the remaining 16,001 trees were used to calculate posterior probabilities in the majority rule consensus tree. After the analysis was finished, each run was examined using the program Tracer v1.5 (Drummond and Rambaut, 2007) to determine burn-in and confirm that both runs had converged. ML analyses were conducted with RAxMLGUI (Silvestro and Michalak, 2012). The GTRGAMMA model was used for all partitions, in accordance with recommendations in the RAxML manual against the use of invariant sites. The parameters of the GTR model used to analyze the dataset were estimated based on the following frequencies, A = 0.242448, C = 0.266436, G = 0.262106, and T = 0.229010; substitution rates AC = 1.080009, AG = 1.915942, AT = 1.147141, CG = 0.835034, CT = 5.341508, and GT = 1.000000, as well as the gamma distribution shape parameter α = 0.291522. The selected model for BI analysis was K2P+G4 (LSU) and SYM+I+G4 (SSU+ITS+RPB1+TEF). The phylogenetic trees (Figure 2) constructed using ML and BI analyses were largely congruent and strongly supported in most branches. Most genera were clustered into their independent clade. The new strains were clustered into four independent clades. Simplicillium larvatum (DY101731, DY101732, DY10251, and DY10252) had a close relationship with S. obclavatum, S. hymenopterorum Z.F. Zhang and L. Cai, and S. spumae N. Kondo, H. Iwasaki and Nonaka. S. araneae (DY101811 and DY101812), S. guizhouense (DY1005 and DY10052), and S. coleopterorum (SD05381 and SD05382) had a close relationship with S. lanosoniveum.

The topological structure of the network is consistent with that of the phylogenetic tree (Figure 1) and could be used for species relationship analysis (Huson and Bryant, 2006). However, a reticular structure was formed in the phylogenetic network by the split of information conflict or fuzzy signals. Three groups were present in the phylogenetic network (Figure 3).

Group I: New strains DY10051, DY10052, DY101812, SD05381, and SD05382 grouped with Simplicillium cicadellidae (GY11011 and GY11012), S. cylindrosporum Nonaka, Kaifuchi and Masuma (JCM 18169, JCM 18170, JCM 18171, JCM 18172, JCM 18173, JCM 18174, and JCM 18175), S. hymenopterorum (DY101691 and DY101692), S. lanosoniveum (CBS 101267, CBS 123.42, and CBS 704.86), S. lepidopterorum W.H. Chen et al. (GY29132), S. minutense Nonaka, Kaifuchi and Masuma (JCM 18176, JCM 18177, and JCM 18178), S. neolepidopterorum (DY101751 and DY101752), and S. scarabaeoidae (DY101391 and DY101392).

Group II: S. aogashimaense Nonaka, Kaifuchi and Masuma (JCM 18167 and JCM 18168) grouped with S. formicae D.P. Wei and K.D. Hyde (MFLUCC 18–1379), S. hymenopterorum (CGMCC 3.19573), S. obclavatum (CBS 311.74 and JCM 18179), and S. spumae (JCM 39050, JCM 39051 and JCM 39054). The new strains DY10251, DY10252, DY101731 and DY101732 are clustered with S. coccinellidae (DY101791 and DY101792) into an independent subgroup.

Group III: S. album Z.F. Zhang and L. Cai (CGMCC 3.19635) grouped with S. calcicole Z.F. Zhang, F. Liu and L.
Phylogenetic identification of the new strains in the genus *Simplicillium* based on ITS sequence. Statistical support values ($\geq 50\%$) are shown at the nodes for ML bootstrap support/BI posterior probabilities.

Cai (LC5371 and LC5586), *S. formicidae* W.H. Chen et al. (DL10041 and DL10042), *S. lamellicola* (CBS 116.25, KYK00006, UAMH 2055 and UAMH 4785), *S. niveum* Mongkols., Noisrip., and Luangs-ard (BCC 83036), *S. pechmerlense* J. Leplat (CBS 147188), and *S. sympodiophorum* Nonaka, Kaifuchi and Masuma (JCM 18184).
FIGURE 2
Phylogenetic analysis to establish the new species in the genus *Simplicillium* by SSU, ITS, LSU, RPB1, and TEF sequences. Statistical support values (≥ 0.5/50%) are shown at the nodes for ML bootstrap support/BI posterior probabilities.
Reconstruction of the neighbor-net network based on ITS sequences from taxa in Figure 1.

Taxonomy

**Simplicillium araneae** W.H. Chen, Y.F. Han, J.D. Liang and Z.Q. Liang, sp. nov.

**Mycobank:** 844146.

**Type:** CHINA, Guizhou, Qiannan Buyi, and Miao Autonomous Prefecture, Duyun City (26°21’27.96”N, 107°22’48.22”E). On a dead spider (Araneae), 1 October 2019, Wanhao Chen, GZAC DY10181 (holotype), ex-type living cultures, DY101811.

**Description:** The colonies showed moderate growth on PDA, reaching a diameter of 31–33 mm in 14 days at 25°C, were convex, with white velutinate aerial mycelium on the front and an yellowish to brown mycelium on the reverse, especially in the middle and entire margins, and soluble pigment was not produced. Vegetative hyphae were branched, hyaline, smooth-walled, septate, and 1.2–1.8 µm wide. The phialides produced on the aerial hyphae were always solitary, aseptate, hyaline, smooth-walled, relatively slender, tapering toward the tip, and 32.9–47.1 × 1.2–2.4 µm in size. Conidia hyaline was ellipsoidal to globose, aseptate, smooth-walled, 1-celled, and 1.8–2.9 × 1.2–1.8 µm in size. Octahedral crystals were absent, and a sexual state was not observed.

**Etymology.** Referring to the ability to colonize spiders.

Additional strain examined. China, Guizhou, Qiannan Buyi, and Miao Autonomous Prefecture, Duyun City (26°21’27.96”N, 107°22’48.22”E). On a dead spider (Araneae), 1 October 2019, Wanhao Chen, DY101812.

Notes: *Simplicillium araneae* was identified as belonging to *Simplicillium* because of its solitary phialides (Figure 4) and the analysis ITS sequence (Figure 1). Compared to the typical characteristics of 23 species, *S. araneae* is morphologically similar to *S. formicae*, *S. hymenopterorum*, and *S. neolepidopterorum* based on the absence of a slime head and octahedral crystals. However, based on the combined dataset of SSU, ITS, LSU, RPB1, and TEF sequences (Figure 2), *S. araneae* clustered into an independent clade and was distinguished from other *Simplicillium* species.

**Simplicillium coleopterorum** W.H. Chen, Y.F. Han, J.D. Liang, and Z.Q. Liang, sp. nov.

**Mycobank:** 844147.

**Type:** CHINA, Guizhou, Qiannan Buyi, and Miao Autonomous Prefecture, Sandu County (25°57’22.21” N, 107°57’54.69” E). On a beetle (Coleoptera), 1 May 2019, Wanhao Chen, GZAC SD0538 (holotype), ex-type living cultures, SD05381.

**Description:** The colonies showed moderate growth on PDA, reaching a diameter of 49–50 mm in 14 days at 25°C, and convex, with white velutinate aerial mycelium on the front and pale brown to brown mycelium on the reverse, especially in the middle and entire margins, and soluble pigment was not produced. The vegetative hyphae were branched, hyaline, smooth-walled, septate, and 1.0–1.6 µm wide. The phialides produced on the aerial hyphae were always solitary, aseptate, hyaline, smooth-walled, relatively slender, tapering toward the...
Simplicillium araneae (A) Infected spider (Araneae); (B, C) PDA-containing culture plate showing the front (B) and reverse (C) sides of the colony; (D–K) solitary phialides and conidia. Scale bars: 10 mm (B, C) and 10 µm (D–K).

Notes:

Simplicillium coleopterorum was identified as belonging to Simplicillium because of its solitary phialides, conidia adhering in subglobose slimy heads, and the absence of octahedral crystals (Figure 5), supported by phylogenetic analysis of ITS sequence (Figure 1). Compared with the typical characteristics of 23 species, S. coleopterorum was morphologically similar to S. cicadellidae, S. coccinellidae, S. formicidae, S. lepidopterorum, S. niveum, S. scarabaeoidae, and S. yunnanense (Figure 6). However, based on the combined dataset of SSU, ITS, LSU, RPB1, and TEF sequences (Figure 2), S. coleopterorum was clustered into an independent clade and distinguished from S. cicadellidae, S. coccinellidae, S. formicidae, S. lepidopterorum, S. niveum, S. scarabaeoidae, and S. yunnanense.

Simplicillium guizhouense W.H. Chen, Y.F. Han, J.D. Liang and Z.Q. Liang sp. nov.

Mycobank: 844148.

Type: CHINA, Guizhou, Qiannan Buyi and Miao Autonomous Prefecture, Duyun City (26°21’27.96”N, 107°22’48.22”E). On an ant (Formicidae), 1 October 2019, Wanhao Chen, GZAC DY1005 (holotype), ex-type living cultures, DY10051.

Description: The colonies showed moderate growth on PDA, reaching a diameter of 35–36 mm in 14 days at 25°C, and were convex, with white velutinate aerial mycelium on the front and yellowish to pale yellowish mycelium on the reverse, especially in the middle and entire margin, and soluble pigment was not produced. The vegetative hyphae were branched, hyaline,
Simplicillium coleopterorum (A) Infected insect (Coleoptera); (B,C) PDA-containing culture plate showing the front (B) and reverse (C) sides of the colony; (D–K) solitary phialides and conidia. Scale bars: 10 mm (B,C) and 10 µm (D–K).

smooth-walled, septate, and 1.4–1.5 µm wide. The phialides produced on the aerial hyphae were always solitary, aseptate, hyaline, smooth-walled, relatively slender, and tapering toward the tip, and 21.1–52.2 × 1.0–1.8 µm in size. Conidia were observed as small globose slimy heads at the apex of the phialides, hyaline, ellipsoidal in shape, aseptate, smooth-walled and 1-celled, and 2.4–2.9 × 1.6–1.8 µm in size. Octahedral crystals were absent, and a sexual state was not observed.

Etymology: Referring to the place where the fungus was collected.

Additional strain examined: China, Guizhou, Qiannan Buyi and Miao Autonomous Prefecture, Duyun City (26°21′27.96″N, 107°22′48.22″E). On an ant (Formicidae), 1 October 2019, Wanhao Chen, DY10052.

Notes: Simplicillium guizhouense is morphologically similar to S. cicadellidae, S. coccinellidae, S. formicidae, S. lepidopterorum, S. niveum, S. scarabaeoidea, and S. yunnanense (Figure 6). Based on the combined dataset of SSU, ITS, LSU, RPB1, and TEF sequences (Figure 2), S. guizhouense clustered into an independent clade, and was distinguished from S. cicadellidae, S. coccinellidae, S. formicidae, S. lepidopterorum, S. niveum, S. scarabaeoidea, and S. yunnanense.

Simplicillium larvatum W.H. Chen, Y.F. Han, J.D. Liang and Z.Q. Liang, sp. nov.

Mycobank: 844149.

Type: CHINA, Guizhou, Qiannan Buyi and Miao Autonomous Prefecture, Duyun City (26°21′27.96″N, 107°22′48.22″E). On a larva (Lepidoptera), 1 October 2019, Wanhao Chen, GZAC DY10173 (holotype), ex-type living cultures, DY101731.

Description: The colonies showed moderate growth on PDA, reaching a diameter of 40–43 mm in 14 days at 25°C, and were convex, with white velutinate aerial mycelium on the front and yellowish mycelium on the reverse, especially in the middle and entire margin, and soluble pigment was not produced. The vegetative hyphae were branched, hyaline, smooth-walled, septate, and 0.9–1.6 µm wide. The phialides produced on the aerial hyphae were always solitary, aseptate, hyaline, smooth-walled, relatively slender, tapering toward the tip, and 16.4–28.7 × 1.2–1.7 µm in size. Conidia were observed as small sub-globular slimy heads at the apex of the phialides, hyaline, ellipsoidal to long ellipsoidal in shape, aseptate, smooth-walled and 1-celled, and 1.8–3.3 × 1.6–2.0 µm
in size. Octahedral crystals were absent, and a sexual state was not observed.

Etymology: Referring to its insect host, a larva.

Additional strain and specimen examined. China, Guizhou, Qiannan Buyi and Miao Autonomous Prefecture, Duyun City (26°21′27.96″N, 107°22′48.22″E). On a larva (Lepidoptera), 1 October 2019, Wanhao Chen, DY101732; on a pupa (Lepidoptera), 1 October 2019, Wanhao Chen, GZAC DY1025, living cultures, DY10251, DY10252.

Notes: *Simplicillium larvatum* is morphologically similar to *S. cicadellidae*, *S. coccinellidae*, *S. lepidopterorum*, *S. niveum*, *S. scarabaeoidea*, and *S. yunnanense* (Figure 7). However, based on the combined dataset of SSU, ITS, LSU, RPB1, and TEF sequences (Figure 2), *S. larvatum* was clustered into an independent clade and phylogenetically close to *S. humicola*, *S. obclavatum*, and *S. spumae*. However, *S. larvatum* was morphologically distinguished from *S. humicola*, which has bigger conidia (3.0–5.0 × 1.5–3.0 μm) with octahedral crystals present. *S. larvatum* was morphologically distinguished from *S. obclavatum*, which has longer phialide (30–52 × 0.8–2.0 μm) with octahedral crystals present. *S. larvatum* was morphologically distinguished from *S. spumae*, which has subglobose or oval to ellipsoidal and octahedral crystals present.

**Discussion**

Zare and Gams (2001) noted that *Simplicillium* species are widely distributed and commonly found on various substrates or as fungicolous fungi. *S. album*, *S. calcicola*, *S. cylindrosporum*, *S. humicola*, *S. minatense*, *S. obclavatum*, *S. pechmerlense*, *S. subtropicum*, and *S. symphodesporophorum* were isolated from soil, marine water, rock, decaying wood, and air (Zare and Gams, 2001; Liu and Cai, 2012; Nonaka et al., 2013; Liang et al., 2017; Zhang et al., 2020b; Leplat et al., 2021). *S. aogashimaense* was isolated from the soil and has also been reported as a symbiotic fungus (Nonaka et al., 2013; Shentu et al., 2020). *S. lansonivieveum* was reported as both an endophytic and
FIGURE 7

Simplicillium larvatum (A) Infected larva (Lepidoptera); (B,C) PDA-containing culture plate showing the front (B) and reverse (C) sides of the colony; (D–L) solitary phialides and conidia. Scale bars: 10 mm (B,C) and 10 µm (D–L).

In the present study, the phylogenetic network was reconstructed to explore the evolutionary relationship, consistent with the phylogenetic tree of the ITS sequence and the combined datasets (SSU, ITS, LSU, RPB1, TEF). According to the phylogenetic network (Figure 3), Simplicillium guizhouense (DY10051 and DY10052) may share a common ancestor with Simplicillium araneae (DY101811 and DY101812), Simplicillium lanosonivum (CBS 123.42 and CBS 704.86), and Simplicillium neolepidopterorum (DY101751 and DY101752). Simplicillium lepidopterorum (GY29131 and GY29132) may share a common ancestor with Simplicillium minatense (JCM 18176, JCM 18177, and JCM 18178). Simplicillium obclavatum (CBS 311.74 and JCM 18179) may share a common ancestor with Simplicillium formicace
(MFLUCC 18–1379), *S. humicola* (CGMCC 3.19573), and *S. spumae* (JCM 39050, JCM 39051 and JCM 39054). *S. calcicole* (LC5371 and LC5586) may share a common ancestor with *S. lanicicola* (CBS 116.25, UAMH 2055, KYK00006, UAMH 4785).

Host shift is usually described as an evolutional process between fungi and their hosts and is often determined by nutrient requirements (Vega et al., 2009). The nutritional model of Hypocreales fungi goes from plants (including living plants and plant residues) to animals (especially insects) and finally to fungi (Spatafora et al., 2007). The results of the phylogenetic network suggest that *S. araneae*, *S. lanosoniveum*, and *S. neolepidopterorum* may have originally come from insects and then jumped to a spider host, plant and fungi substrate, or another insect host, respectively. *S. lepidopterorum* may have originally come from the soil and then jumped to an insect host. *S. formicae* and *S. spumae* may have originally come from air or soil and then jumped as hyperparasitic fungi or water substrates. *S. lanicicola* may have originally come from rock substrate and then jumped as hyperparasitic fungus. These results suggest that host jump may be common in *Simplicillium* species.

*S. coleopterorum* and *S. larvatum* could not split from *S. scarabaeoidea* and *S. coccinellidae* in the phylogenetic network as only the ITS sequence was analyzed. However, they could be phylogenetically distinguished by a multi-locus phylogenetic analysis. Therefore, more sequence information should be added to the phylogenetic network in order to analyze their evolutionary relationship. Moreover, *S. lanosoniveum* was transferred to the genus *Simplicillium* by Zare and Gams (2001) with the strain CBS 123.42. In the present study, three strains of *S. lanosoniveum* (CBS 101267, CBS 123.42, and CBS 704.86) were tested. Strains CBS 123.42 and CBS 704.86 were clustered into a subclade. However, strain CBS 101267 was clustered with four strains of *S. subtropicum* (JCM 18180, JCM 18181, JCM18182, and JCM 18183). The pairwise dissimilarity of ITS sequences shows only a 4 bp difference, with 552 bp between strains CBS 101267 and JCM18180. This result supports strain CBS 101267 being identified as *S. subtropicum*.

**Author contributions**

WC isolated the fungi, built up the phylogenetic tree, and wrote the manuscript. YH identified the fungal isolates, revised the manuscript, and provided partial funding. JL, XR, and JZ revised the manuscript and provided partial funding. ZL identified the fungal isolates and revised the manuscript. All authors discussed the results.

**Funding**

This work was supported by the National Natural Science Foundation of China (Grant No. 31860002), the High-level Innovative Talents Training Object in Guizhou Province (No. Qiankehepintagiairencai [2020]6005), the Science and Technology Foundation of Guizhou Province (No. Qiankehejichu [2020]1Y060), the Program of Innovative Scientific and Technological Talent Team of Guizhou Province (2020-5010), and the Construction Program of Guizhou Engineering Research Center (Qian Fa Gai Gao Ji 2020-896).

**Conflict of interest**

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

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**Data availability statement**

The original contributions presented in the study are included in the article /Supplementary material, further inquiries can be directed to the corresponding author.

**Supplementary material**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2022.950773/full#supplementary-material

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**Supplementary material**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2022.950773/full#supplementary-material
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