Morphology and phylogeny reveal two novel *Coryneum* species from China

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Abstract

*Coryneum* is currently the sole genus of Coryneaceae in Diaporthales, distinguished from other diaporthalean genera by transversely distoseptate brown conidia. However, *Coryneum* species are presently difficult to identify because of variability and overlap of morphological characters and the lack of sequence data for most described species. During fungal collection trips in China, 13 *Coryneum* isolates were obtained from cankered branches of *Ilex* and *Quercus*. Morphological and phylogenetic analyses (ITS, LSU, TEF1-α and RPB2) revealed that these strains belong to two new species (viz. *Coryneum ilicis* sp. nov. and *C. songshanense* sp. nov.), and three known species, *C. gigasporum*, *C. sinense*, and *C. suttonii*. *Coryneum ilicis* has larger conidia and more distosepta than most *Coryneum* species. *Coryneum songshanense* was similar to *C. sinense* from the same host genus, *Quercus*, in conidial length, but distinct in conidial width and by molecular data.

Keywords

*Coryneaceae*, Diaporthales, systematics, taxonomy

Introduction

The genus *Coryneum* Nees is currently the only accepted genus in Coryneaceae and it forms a distinct phylogenetic lineage in Diaporthales (Senanayake et al. 2017, 2018, Voglmayr et al. 2017, Fan et al. 2018a, Jiang et al. 2018, Senwanna et al. 2018, Wijayawardene et al. 2017, 2018). The genus *Coryneum* was introduced based on the asexual
morph, with *C. umbonatum* Nees as the type species (Nees von Esenbeck 1816), and the sexual morph *Pseudovalsa* Ces. & De Not. was introduced later, based on *P. lanciformis* (Fr.) Ces. & De Not. (Cesati & De Notaris 1863). *Coryneum* was recommended to be adopted due to priority and the need of fewer new combinations (Rossman et al. 2015). Most *Coryneum* species were considered as phytopathogens, which were discovered from cankers and dieback of shoots and twigs (Wijayawardene et al. 2016, Senanayake et al. 2017, Jiang et al. 2018). However, diseases are commonly mild and only rarely cause serious symptoms in the hosts. Additionally, pathogenicity tests have not yet been conducted.

*Coryneum* species are generally considered highly host-specific, and 28 species and a variety were accepted in this genus before this study (Sutton 1975, 1980, Wijayawardene et al. 2016, Jiang et al. 2018, Senwanna et al. 2018). *Coryneum terrophilum* was the only species isolated from soil, and the others were reported from dead branches (Table 1). Fagales species are the major hosts of *Coryneum* species, and host trees from other orders are also hardwoods with rough barks (Table 1).

Molecular phylogenies based on multi-gene loci including the internal transcribed spacer (ITS) and the large subunit (LSU) regions of the nuclear rDNA, translation elongation factor-1α (*TEF1*-α) and the second largest subunit of the RNA polymerase II (*RPB2*) have been widely used to infer species delimitation within many genera in Diaporthales (Voglmayr et al. 2012, 2017, 2019, Voglmayr and Jaklitsch 2014, Fan et al. 2018b, Jiang et al. 2019), and are particularly important in speciose genera like *Coryneum*. Hence, DNA extraction from known species and fresh collections from the potential hosts will greatly improve the elucidation of species concept and circumscription in *Coryneum*. Thus, the main objectives of the present study were to identify *Coryneum* taxa based on morphology and phylogenetic evidence, and to analyse the relationships between *Coryneum* species and host genera.

**Materials and methods**

**Sample collection and isolation**

Sample collection trips were conducted in Beijing, Hebei and Shaanxi Provinces of China during June to October in 2017 and 2018, aiming to collect fresh specimens with *Coryneum*-like taxa. Fagales plants were the main hosts and other hardwoods with rough barks were also investigated. Healthy branches and twigs were covered by green leaves, hence the dying and dead materials were conspicuous during our investigations. Asexual fruiting bodies were easily discovered as black spots on the host barks. Tree tissues with fruiting bodies were cut into small pieces, packed in paper bags and taken to the laboratory for further studies. Isolations were obtained by removing the ascospores or conidial masses from the fruiting bodies on to clean potato dextrose agar (PDA) plates, which were incubated at 25 °C until spores germinated. Single germinating spores were transferred on to new PDA plates, which were kept at 25 °C in the dark. Specimens were deposited at the Museum of the Beijing Forestry University (BJFC) and axenic cultures are maintained at the China Forestry Culture Collection Centre (CFCC).
Table 1. Hosts, conidial sizes, and numbers of distosepta of currently accepted Coryneum species.

| Species            | Host genus | Host family | Host order | Conidial size (μm) | No. of distosepta | References                  |
|--------------------|------------|-------------|------------|--------------------|-------------------|-----------------------------|
| C. arausiacum      | Quercus    | Fagaceae    | Fagales    | 42–56 × 13–16      | 4–5               | Senanayake et al. (2017)   |
| C. betulinum       | Betula     | Betulaceae  | Fagales    | 31–36 × 14–17      | 4–5               | Sutton (1975)               |
| C. calophylli      | Calophyllum| Guttiferae  | Parietales | 38–48 × 12.5–14.5  | 5–6               | Sutton (1975)               |
| C. carpiniola      | Carpinus   | Betulaceae  | Fagales    | 50–68 × 8–11       | 7–11              | Sutton (1975)               |
| C. castaneicola    | Castanea   | Fagaceae    | Fagales    | 56–80 × 9.5–13     | 5–8               | Sutton (1975)               |
| C. ceatti          | Aesculus   | Hippocastanaceae | Sapindales | 80–90 × 13–15     | 6–7               | Sutton (1975)               |
| C. clusiae         | Clusia     | Clusiaceae  | Malpighiales | 30–40 × 20–30 | 3–5               | Sutton (1975)               |
| C. compactum       | Ulmus      | Ulmaceae    | Urticales  | 40–58 × 15–21      | 4–6               | Sutton (1975)               |
| C. depressum       | Quercus    | Fagaceae    | Fagales    | 44–53 × 19–23      | 4–6               | Sutton (1975)               |
| C. elevatum        | Quercus    | Fagaceae    | Fagales    | 56–69 × 24–28      | 5–7               | Sutton (1975)               |
| C. gigaporum       | Castanea   | Fagaceae    | Fagales    | 88–117 × 18–23     | 7–9               | Jiang et al. (2018)         |
| C. gregoryi        | Eucalyptus | Myrtaceae   | Myrtiales  | 32.5–43 × 12–16    | 5–9               | Sutton and Sharma (1983)    |
| C. heveanum        | Hevea      | Euphorbiaceae | Malpighiales | 40–68 × 14–20 | 4–6               | Senwanna et al. (2018)      |
| C. ilicis          | Ilex       | Aquifoliaceae | Sapindales | 82–105 × 9.5–12.5  | 10–11             | This study                  |
| C. japonicum       | Quercus    | Fagaceae    | Fagales    | 45–60 × 11–12      | 5–7               | Sutton (1975)               |
| C. lanciforme      | Betula     | Betulaceae  | Fagales    | 45–53 × 16–18      | 4–6               | Sutton (1975)               |
| C. megaspermum     | Quercus    | Fagaceae    | Fagales    | 73–97 × 13–16      | 7–11              | Sutton (1980)               |
| C. megaspermum var. cylindricum | Quercus | Fagaceae | Fagales | 100–125 × 10–13 | 7–8               | Sutton (1975)               |
| C. modonatum       | Castanea   | Fagaceae    | Fagales    | 50–71 × 14–19      | 5–8               | Sutton (1975)               |
| C. neesi           | Quercus    | Fagaceae    | Fagales    | 68–82 × 18–22      | 6–8               | Sutton (1975)               |
| C. pruni           | Prunus     | Rosaceae    | Rosales    | 14–23 × 5.5–9      | 4–5               | Wijayawardene et al. (2016) |
| C. psidi           | Psidium    | Myrtaceae   | Myrtiales  | 25–40 × 14–17      | 5–6               | Sutton (1975)               |
| C. pyricola        | Pyrus      | Rosaceae    | Rosales    | 61–70 × 24–32      | 5–7               | Sutton (1975)               |
| C. quercinum       | Quercus    | Fagaceae    | Fagales    | 45–60 × 14–16      | 6–7               | Mushumary and Sutton (1986) |
| C. sinense         | Quercus    | Fagaceae    | Fagales    | 50–76 × 13–17      | 5–7               | Jiang et al. (2018)         |
| C. songbenense     | Quercus    | Fagaceae    | Fagales    | 51–76 × 9–11.5     | 5–7               | This study                  |
| C. stromatoidesm   | Tsuga      | Pinaceae    | Pinales    | 105–180 × 16–20    | 9–17              | Sutton (1975)               |
| C. suttonii        | Castanea   | Fagaceae    | Fagales    | 60–76 × 10–14.5    | 4–5               | Jiang et al. (2018)         |
| C. sydowianum      | Abris      | Betulaceae  | Fagales    | 50–58 × 14–17      | 5–6               | Sutton (1975)               |
| C. terrophilum     | NA         | NA          | NA         | 25–55 × 15–24      | 3–7               | Sutton and Sharma (1983)    |
| C. umbonatum       | Quercus    | Fagaceae    | Fagales    | 57–72 × 13–16      | 5–7               | Sutton (1975)               |

Morphological analysis

Species identification was based on the morphological characters of the sexual and asexual morphs produced on natural substrates. Cross-sections were prepared manually using a double-edged blade under a Leica stereomicroscope (M205 FA). Photomicrographs were captured with a Nikon Eclipse 80i microscope equipped with a Nikon digital sight DS-Ri2 high-definition colour camera, using differential interference contrast (DIC) illumination and the Nikon software, NIS-Elements D Package 3.00. Measurements of ascospores and conidia are reported as the maximum and minimum in parentheses and the range representing the mean ± standard deviation of the number of measurements is given in parentheses (Voglmayr et al. 2017). Cultural characteristics of isolates incubated on MEA in the dark at 25 °C were recorded.

Recognition and identification of Coryneum species were based on fruiting bodies formed on tree bark, supplied by conidiomata produced on PDA plates. Ascomata and conidiomata from tree bark were sectioned by hand using a double-edged blade,
and conidiomata from PDA plates were picked using a needle, which were observed under a dissecting microscope. At least 10 conidiomata/ascomata, 10 asci, and 50 conidia/ascospores were measured to calculate the mean sizes and standard deviation. Microscopy photographs were captured with a Nikon Eclipse 80i compound microscope equipped with a Nikon digital sight DS-Ri2 high definition colour camera, using differential interference contrast illumination.

**DNA extraction, PCR amplification and sequencing**

Genomic DNA was extracted from colonies grown on cellophane-covered PDA plates using a modified CTAB method (Doyle and Doyle 1990). PCR amplifications were performed in a DNA Engine Peltier Thermal Cycler (PTC-200; Bio-Rad Laboratories, Hercules, CA, USA). The primer sets ITS1/ITS4 (White et al. 1990) were used to amplify the ITS region. The primer pair LR0R/LR5 (Vilgalys and Hester 1990) was used to amplify the LSU region. The primer pairs EF1-688F/EF1-986R or EF1-728F/TEF1-LLErev (Carbone and Kohn 1999, Jaklitsch et al. 2006, Alves et al. 2008) were used to amplify TEF1-a gene. The primer pair dRPB2-5f/dRPB2-7r (Voglmayr et al. 2016) was used to amplify the RPB2 gene. The polymerase chain reaction (PCR) assay was conducted as described by Fan et al. (2018a). PCR amplification products were assayed via electrophoresis in 2 % agarose gels. DNA sequencing was performed using an ABI PRISM® 3730XL DNA Analyzer with a BigDye Terminater Kit v.3.1 (Invitrogen, USA) at the Shanghai Invitrogen Biological Technology Company Limited (Beijing, China). Novel sequences generated in the current study were deposited in GenBank (Table 2).

**Phylogenetic analyses**

Sequences generated from the above primers of the different genomic regions (ITS, LSU, TEF1-a and RPB2) were analysed in comparison to known species, Stilbospora macrosperrna (CBS 115073) and Stegonsporium pyriforme (CBS 120522) were used as the outgroup taxa (Jiang et al. 2018). All sequences were aligned using MAFFT v. 6 (Katoh and Toh 2010) and edited manually using MEGA v. 6 (Tamura et al. 2013). Phylogenetic analyses were performed using PAUP v. 4.0b10 for maximum parsimony (MP) analysis (Swofford 2003), and PhyML v. 3.0 for Maximum Likelihood (ML) analysis (Guindon et al. 2010).

A partition homogeneity test with heuristic search and 1000 replicates was performed using PAUP v. 4.0b10 to assess incongruence among the ITS, LSU, TEF1-a, and RPB2 sequence datasets in reconstructing phylogenetic trees. MP analysis was run using a heuristic search option of 1000 search replicates with random-addition of sequences with a tree bisection and reconnection (TBR) algorithm; branches of zero length were collapsed (collapse = minbrlen), and all equally most parsimonious trees were saved. Other calculated parsimony scores were tree length (TL), consistency index (CI), retention index (RI), and rescaled consistency (RC). ML analysis was performed
Table 2. Strains used in the phylogenetic tree and their culture accession and GenBank numbers. Strains from this study are in bold.

| Species            | Strains      | GenBank numbers |
|--------------------|--------------|-----------------|
|                    |              | ITS            | LSU            | TEF1-α         | RPB2         |
| Coryneum castaneicola | CFCC 52315   | MH683551       | MH683559       | MH685731       | MH685723     |
| Coryneum castaneicola | CFCC 52316   | MH683552       | MH683560       | MH685732       | MH685724     |
| Coryneum depressum  | D202         | MH674330       | MH674330       | MH674338       | MH674334     |
| Coryneum heveanum   | MFLUCC 17-0369 | MH778707       | MH778703       | MH780881       | NA           |
| Coryneum heveanum   | MFLUCC 17-0376 | MH778708       | MH778704       | NA             | NA           |
| Coryneum gigasporum | CFCC 52319   | MH683557       | MH683565       | MH685737       | MH685729     |
| Coryneum gigasporum | CFCC 52320   | MH683558       | MH683566       | MH685738       | MH685730     |
| Coryneum gigasporum | G14          | MK799957       | MK799944       | MK799830       | MK799820     |
| Coryneum gigasporum | G15          | MK799958       | MK799945       | MK799831       | MK799821     |
| Coryneum ilicis     | CFCC 52994   | MK799948       | MK799935       | NA             | NA           |
| Coryneum ilicis     | CFCC 52995   | MK799949       | MK799936       | NA             | NA           |
| Coryneum ilicis     | CFCC 52996   | MK799950       | MK799937       | NA             | NA           |
| Coryneum lanciforme | D215         | MH674332       | MH674332       | MH674340       | MH674336     |
| Coryneum modonium   | D203         | MH674331       | MH674331       | MH674339       | MH674335     |
| Coryneum modonium   | CBS 130.25   | MH854812       | MH866313       | NA             | NA           |
| Coryneum sinense    | CFCC 52452   | MH683553       | MH683561       | MH685733       | MH685725     |
| Coryneum sinense    | CFCC 52453   | MH683554       | MH683562       | MH685734       | MH685726     |
| Coryneum sinense    | X20          | MK799952       | MK799939       | MK799825       | MK799815     |
| Coryneum sinense    | X23          | MK799953       | MK799940       | MK799826       | MK799816     |
| Coryneum sinense    | X60          | MK799951       | MK799938       | MK799824       | MK799814     |
| Coryneum songshanense | CFCC 52997 | MK799946       | MK799933       | MK799822       | MK799812     |
| Coryneum songshanense | CFCC 52998 | MK799947       | MK799934       | MK799823       | MK799813     |
| Coryneum suttonii   | CFCC 52317   | MH683555       | MH683563       | MH685735       | MH685727     |
| Coryneum suttonii   | CFCC 52318   | MH683556       | MH683564       | MH685736       | MH685728     |
| Coryneum suttonii   | Z15-1        | MK799954       | MK799941       | MK799827       | MK799817     |
| Coryneum suttonii   | Z17          | MK799955       | MK799942       | MK799828       | MK799818     |
| Coryneum suttonii   | Z86          | MK799956       | MK799943       | MK799829       | MK799819     |
| Coryneum umbonatum  | D201         | MH674329       | MH674329       | MH674337       | MH674333     |

using a GTR site substitution model, including a gamma-distributed rate heterogeneity and a proportion of invariant sites (Guindon et al. 2010). The branch support was evaluated using a bootstrapping method of 1000 bootstrap replicates (Hillis and Bull 1993). The MP bootstrap analyses were done with the same settings as for the heuristic search, but with 10 rounds of heuristic search during each bootstrap replicate. Phylogenograms were shown using FigTree v. 1.4.3 (Rambaut 2016).

Results

Phylogenetic analyses

The alignment based on the combined sequence dataset (ITS, LSU, TEF1-α, and RPB2) included 30 ingroup taxa and two outgroup taxa (Stilbospora macrosperma and Stegonsporium pyriforme), comprising 3544 characters in the aligned matrix. Of these, 2570 characters were constant, 267 variable characters were parsimony-uninformative and 706 characters were parsimony informative. The partition homogeneity test resulted in an insignificant value (level 95%), indicating that ITS, LSU, TEF1-α and
Figure 1. Phylogenetic tree based on an MP analysis of a combined DNA dataset of ITS, LSU, TEF1-α and RPB2 gene sequences for the species of *Coryneum*. Bootstrap values ≥ 50% for MP/ML analyses are presented at the branches. Scale bar = 50 nucleotide substitutions.

RPB2 sequence dataset could be combined. The MP analysis resulted in 2 equally most parsimonious trees; the first tree (TL = 1624, CI = 0.784, RI = 0.822, RC = 0.645) is shown in Fig. 1. The two MP trees were identical, except for an interchanged position of *C. ilicis* and *C. songshanense* (not shown). Tree topology of the best tree revealed by the ML analyses was identical to that of the MP tree shown. The phylogram based on the four gene sequences showed that the accessions here studied represented 2 new and 3 known species in *Coryneum* (Fig. 1).
Taxonomy

*Coryneum ilicis* C.M. Tian & N. Jiang, sp. nov.
MycoBank: MB830201

Figure 2

**Diagnosis.** *Coryneum ilicis* is characterised by its host, *Ilex pernyi*, and large conidia with 10–11 distosepta.

**Holotype.** CHINA. Shaanxi Province: Zhashui County, on branches of *Ilex pernyi*, 12 August 2017, N. Jiang (holotype: BJFC-S1720; ex-type culture from ascospore: CFCC 52994; living culture from conidium: CFCC 52996).

**Etymology.** Named after the host genus on which it was collected, *Ilex*.

**Description.** Associated with canker on branches of *Ilex pernyi*. *Sexual morph:* Pseudeostromata 0.5–1.5 mm diam., typically distinct, circular, without perithecial bumps, containing 1 or 2 perithecia embedded in a well-developed entostroma. Central column and entostroma grey. Ostioles inconspicuous and often invisible at the surface of the ectostromatic disc. Perithecia (350–)500–700(–850) μm diam. (n = 20), globular, somewhat flattened at the base. Asci 110–155 × 13–20 μm, 8-spored, unitunicate, clavate, shortly pedicellate, apically rounded, with a conspicuous apical ring. Ascospores (26.2–)29.7–35.5(–36.2) × (11.0–)11.8–14.3(–15.2) μm, l/w = (1.9–)2.2–2.9(–3.2) (n = 50), 1-seriate, fusiform, ends pointed, uniseptate, constricted at the septa, hyaline, guttulate, smooth-walled. *Asexual morph:* Conidiomata acervular, 0.2–1 mm wide, 0.2–1.2 mm high, solitary, erumpent through the outer periderm layers of the host, scattered, surface tissues above slightly domed. Conidiophores 40–85 μm long, 3–7 μm wide, branched, cylindrical, septate, hyaline at the apex, pale brown at the base. Conidiogenous cells holoblastic, integrated, indeterminate, cylindrical, expanding towards the apices, pale brown, smooth, with 0–1 percurrent extensions. Conidia (82–)87–95(–105) × (9.5–)10.5–11.5(–12.5) μm, l/w = (7.4–)7.7–9.1(–9.3) (n = 50), variable in shape, curved, broadly fusiform to fusiform, cylindrical or clavate, dark brown, smooth-walled, 10–11-distoseptate, apical cell with a hyaline tip, truncate and black at the base.

**Culture characters.** On PDA at 25 °C, colonies growing slowly and unevenly, reaching 70 mm diam. within 25 d, gradually becoming brownish dark grey in colour with scant cottony aerial mycelium, asexual morphs developed after 35 d.

**Additional specimen examined.** CHINA. Shaanxi Province: Zhashui County, on branches of *Ilex pernyi*, 12 August 2017, N. Jiang (isotype: BJFC-S1721; living culture: CFCC 52995).

**Notes.** *Coryneum ilicis* is the sole species known from the host genus *Ilex*; it can be easily recognised by host association and phylogeny (Fig. 1). Morphologically, conidia of *Coryneum ilicis* are larger and have more distosepta than in most of the other species (Table 1).
Figure 2. *Coryneum ilex* from *Ilex pernyi* (BJFC-S1720, holotype)  

A Fruiting bodies on natural substrate in surface view  

B pseudostroma in transverse section, showing perithecia and gray entostroma  

C longitudinal sections through pseudostromata  

D ascus  

E–J ascospores  

K conidiophores  

L–N conidia. Scale bars: 1 mm (A); 0.5 mm (B, C); 20 μm (D); 10 μm (E–N).
Coryneum songshanense C.M. Tian & N. Jiang, sp. nov.
Mycobank: MB830202
Figure 3

Diagnosis. Coryneum songshanense can be distinguished from the morphologically similar C. sinense by its narrower conidia.

Holotype. CHINA. Beijing City: Songshan Mountain, on dead twigs of Quercus dentata, 15 June 2018, N. Jiang & C.M. Tian (holotype: BJFC-S1722; ex-type culture from ascospore: CFCC 52997).

Etymology. Named after the mountain on which it was collected, Songshan Mountain.

Description. Associated with canker on twigs of Quercus dentata. Sexual morph: Pseudostromata 0.3–1 mm diam., typically distinct, circular, without perithecial bumps, containing up to 6 perithecia embedded in a well-developed entostroma. Ectostromatic disc distinct, circular, black, 0.3–0.5 mm diam. Central column and entostroma grey. Ostioles inconspicuous and often invisible at the surface of the ectostromatic disc. Perithecia (150–)200–450(–550) μm diam. (n = 20), globular, somewhat flattened at the base with black short neck. Asci 75–145 × 17–23 μm, 8-spored, unitunicate, clavate, shortly pedicellate, apically rounded, with an inconspicuous apical ring. Ascospores (24.1–)25.5–35.4(–38.2) × (7.5–)7.9–9.8(–10.6) μm, l/w = (3.0–)3.3–3.8(–4.2) (n = 50), 2-seriate, fusiform, ends pointed, uniseptate or aseptate, not constricted at the septa, hyaline, guttulate, smooth-walled. Asexual morph: Conidiomata acervular, 0.2–0.6 mm wide, 0.2–0.5 mm high, solitary, erumpent through the outer periderm layers of the host, scattered, surface tissues above slightly domed. Conidiophores 15–35 μm long, 4–7 μm wide, unbranched, cylindrical, septate, hyaline at the apex, pale brown at the base. Conidiogenous cells holoblastic, integrated, indeterminate, cylindrical, expanding towards the apices, pale brown, smooth, with 0–1 percurrent extensions. Conidia (51–)56–67(–76) × (9–)10–11(–11.5) μm, l/w = (5.2–)5.5–6.9(–8.1) (n = 50), variable in shape, curved, broadly fusiform to fusiform, cylindrical or clavate, dark brown, smooth-walled, 5–7-distoseptate, apical cell with a hyaline tip, truncate and black at the base.

Culture characters. On PDA at 25 °C, colonies growing slowly and unevenly, reaching 70 mm diam. within 30 d, gradually becoming brownish dark grey in colour with scant cottony aerial mycelium, asexual morphs developed after 40 d.

Additional specimen examined. CHINA. Beijing City: Songshan Mountain, on dead twigs of Quercus dentata, 15 June 2018, N. Jiang & C.M. Tian (isotype: BJFC-S1723; living culture from conidium: CFCC 52998).

Notes. So far, ten species and one variety have been described from Quercus branches, and they can be distinguished by conidial characteristics (Muthumary and Sutton 1986, Jiang et al. 2018, Table 1). Coryneum songshanense and C. sinense can be distinguished from C. arausiacum, C. depressum, C. elevatum, C. japonicum, C. megaspernum, C. megaspernum var. cylindricum, C. neesii, C. umbonatum, and C. quercinum by unbranched conidiophores (Sutton 1975, Muthumary and Sutton 1986, Jiang
Figure 3. *Coryneum songshanense* from *Quercus dentata* (BJFC-S1722, holotype) A, B Fruiting bodies on natural substrate in surface view C pseudostroma in transverse section, showing perithecia and gray entostroma D longitudinal sections through pseudostromata E, F immature asci G, H immature Ascospores I, J conidiophores K–M conidia. Scale bars: 1 mm (A, B); 0.5 mm (C, D); 10 μm (E–M).
Coryneum species from China

et al. 2018). *Coryneum songshanense* is obviously distinguished from *C. sinense* in narrower conidia (9–11.5 μm in *Coryneum songshanense* vs. 13–17 μm in *C. sinense*) and phylogeny (Fig. 1).

**Discussion**

In this study, fresh *Coryneum* specimens were collected in China and identified based on combined morphological and molecular data. Additional accessions of three recently described *Coryneum* species, *C. gigasporum*, *C. sinense*, and *C. suttonii* (Jiang et al. 2018), were identified, with matching conidial characteristics and sequences (Fig. 1). The new species *C. ilicis* was discovered on *Ilex pernyi* (Aquifoliaceae, Sapindales), which represents a new host family and genus for *Coryneum*. *Coryneum cesatii* was reported from the same host order, Sapindales, on branches of *Aesculus* (Hippocastanaceae) (Sutton 1975). The second new species, *Coryneum songshanense*, was discovered on dead twigs of *Quercus dentata* (Fagaceae, Fagales). Host species belonging to Fagales show higher diversity of *Coryneum* species (Table 1), and it is likely that additional taxa will be discovered by molecular data, considering that in many regions suitable hosts have not yet been adequately studied.

However, most of the *Coryneum* species are lacking DNA sequences, thus species identification based on DNA sequence analyses is presently difficult. Hence, polyphasic approach, i.e. incorporating morphological characters (such as conidial sizes and numbers of distosepta), as well as host associations are important for species identification (Sutton 1975, 1980, Jiang et al. 2018). However, host identifications may be incorrect and many geographical areas remain insufficiently studied. In addition, the morphological characters often significantly overlap between species, which makes identifications solely by morphology challenging. Hence, studies based on the types of already described species and new collections from potential hosts are important to achieve a reliable species classification and circumscription within *Coryneum*.

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