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Convergent evolution of highly reduced fruiting bodies in Pezizomycotina suggests key adaptations to the bee habitat

Anja Amtoft Wynns

Abstract

Background: Among the understudied fungi found in nature are those living in close association with social and solitary bees. The bee-specialist genera Bettsia, Ascosphaera and Eremascus are remarkable not only for their specialized niche but also for their simple fruiting bodies or ascocarps, which are morphologically anomalous in Pezizomycotina. Bettsia and Ascosphaera are characterized by a unicellular cyst-like cleistothecium known as a spore cyst, while Eremascus is characterized by completely naked asci, or asci not formed within a protective ascocarp. Before molecular phylogenetics the placement of these genera within Pezizomycotina remained tentative; morphological characters were misleading because they do not produce multicellular ascocarps, a defining character of Pezizomycotina. Because of their unique fruiting bodies, the close relationship of these bee-specialist fungi and their monophyly appeared certain. However, recent molecular studies have shown that Bettsia is not closely related to Ascosphaera.

In this study, I isolated the very rare fungus Eremascus fertilis (Ascomycota, Pezizomycotina) from the bee bread of honey bees. These isolates represent the second report of E. fertilis both in nature and in the honey bee hive. To establish the systematic position of E. fertilis and Bettsia alvei, I performed phylogenetic analyses of nuclear ribosomal LSU + SSU DNA sequences from these species and 63 additional ascomycetes.

Results: The phylogenetic analyses revealed that Eremascus is not monophyletic. Eremascus albus is closely related to Ascosphaera in Eurotiomycetes while E. fertilis belongs in Myxotrichaceae, a putative member of Leotiomycetes. Bettsia is not closely related to Ascosphaera and like E. fertilis apparently belongs in Leotiomycetes. These results indicate that both the naked ascus and spore cyst evolved twice in the Pezizomycotina and in distantly related lineages. The new genus Skoua is described to accommodate E. fertilis.

Conclusions: The naked ascus and spore cyst are both shown to have evolved convergently within the bee habitat. The convergent evolution of these unusual ascocarps is hypothesized to be adaptive for bee-mediated dispersal. Elucidating the dispersal strategies of these fungal symbionts contributes to our understanding of their interaction with bees and provides insight into the factors which potentially drive the evolution of reduced ascocarps in Pezizomycotina.

Keywords: Ascosphaerales, Eremascaceae, Honey bees, Solitary bees, Yeasts, Skoua

Background

The Ascomycota are an ecologically diverse group of fungi characterized by the production of meiospores in sac-like structures called asci. In the subphylum Pezizomycotina, asci are formed within a protective ascocarp, while in the basal lineages (Saccharomycotina and nearly all members of Taphrinomycotina) asci lack a protective covering and are called naked asci. Four major multicellular ascocarp types are recognized in the Ascomycota: apothecia (cup-shaped ascocarps with an exposed hymenium), perithecia (flask-shaped ascocarps with a pore through which ascospores are released), pseudothecia (ascocarps with asci contained in numerous locules), and cleistothecia (entirely closed ascocarps with no predefined opening and no regular arrangement of asci) [1, 2]. The traditional classification system of Pezizomycotina (Ascomycota) placed great emphasis on these ascocarp types. Subsequently, DNA sequence-based phylogenies have
shown that similar ascocarp types have evolved multiple times in distantly related lineages [3, 4]. As more sequence data become available, it is increasingly clear that morphological convergence of ascocarp types is not uncommon, and that the evolution of ascocarps and modes of ascus and ascocarp dehiscence are frequently associated with common spore dispersal strategies [5, 6, 4].

Phylogenetic studies have identified the major lineages shaping the backbone of the fungal tree of life [7, 8, 4] but the placement of many taxa at and below the class level remains uncertain. Increased DNA sampling of understudied groups, especially those whose systematic placement is based on morphology, may be critical for enabling ancestral-state reconstructions of characteristics such as ascus-dehiscence type, ascocarp ontogeny, and lifestyle and spore dispersal strategies.

The three bee-specialist genera *Ascosphaera*, *Arrhenosphaera* (Ascosphaeraceae; Eurotiomycetes) and *Bettsia* (incertae cedis; formerly Ascosphaeraceae) are among the understudied and seldom-collected fungi. Within the bee habitat these fungi lead both saprotrophic and pathogenic lifestyles. They are found growing on pollen provisions, larval feces (Fig. 1e–f), cocoons, materials used by the bees to construct brood cells, and within bee larvae [9–15]. At least half of the species of *Ascosphaera* are bee brood pathogens infecting the larval stage and causing the bee disease commonly known as chalkbrood. In addition to sharing a specialized ecological niche [16], *Ascosphaera*, *Arrhenosphaera* and *Bettsia* are remarkable for their ascocarps (Figs. 1–2) which are morphologically anomalous in Pezizomycotina. Before DNA sequence data were available, the placement of Ascosphaeraceae among the Fungi and their systematic affinity of the monotypic genus *Arrhenosphaera* to Ascosphaeraceae is based on the shared character of a spore cyst. Unfortunately, no holotype was designated in the protologue of *Arrhenosphaera cranoi* Stejskal [9] and no additional collections or reports of the fungus have been made since its description as a problematic pathogen of honey bees in Venezuela in 1974.

The placement of Ascosphaeraceae in Eurotiales (Eurotiomycetidae) is based on similar early sexual development in *Monascus* and *Aspergillus* [17, 25], and is supported by DNA sequence-based phylogenies [3, 24]. These phylogenies also revealed a close relationship between Ascosphaeraceae and Eremascaceae, a family of just one genus, *Eremascus*. Ascosphaeraceae and *Eremascus* both lack hyphal fruiting bodies and for this reason were loosely referred to as ‘yeasts’ [26]. The shared simple ascocarp morphology of these two taxa was recognized before molecular data became available but their possible relatedness was doubted because of differences in their sexual systems [25, 27].

The genus *Eremascus* includes two species, *E. fertilis* Stoppel and *E. albus* Eidam, and is characterized by naked asci and a predilection for high sugar substrates [28]. Naked asci, with no interspersed sterile hyphae, occur elsewhere only within the early diverging lineages of the Ascomycota: the yeast-like Taphrinomycotina and the true-yeasts, Saccharomycotina [29]. The morphological similarity of *Eremascus* to the yeast-like fungi led some authors to believe that *Eremascus* should be placed among the Saccharomycetales, noting that the genus differs from other members of this group only in lacking asexual reproduction by budding or otherwise [30, 19]. Despite its unique fruiting body type within the Pezizomycotina, *Eremascus* has received little attention in phylogenetic studies. The prevailing view has been that *Eremascus* is monophyletic [31, 28], and its closest relative is Ascosphaeraceae [24, 32, 26]. The supposed monophyly of the genus is based on the shared character of completely naked asci, while the two species (*E. fertilis* and *E. albus*) differ in ascospore morphology and sexual development [27, 33].

Both *Eremascus* species are xerophiles thriving in conditions where water activity is low and solute concentrations are high [28]. The natural habitat of *Eremascus* remains elusive; both species are very rarely collected and have previously been isolated only from various
high-sugar foods such as prunes, preserved fruit, shortcake, plum jam, mincemeat, honeycomb and on pollen [28, 34, 35]. In the process of studying the diversity of spore cyst fungi in the nests of solitary bees and in the hives of honey bees in Denmark, Eremascus fertilis was serendipitously isolated from the bee bread of honey bee hives. This is the second report of E. fertilis in nature: the first report was in 1912 when A. Betts also observed the fungus growing on pollen in honey bee hives [35].

Eremascus fertilis has not previously been included in phylogenetic studies at the class level. In this study, I sequenced the large subunit (LSU) and small subunit (SSU) nuclear ribosomal DNA regions for E. fertilis and also for Bettsia alvei. These sequences were added to a large matrix of sequences from other Ascomycota, which was then analyzed both by maximum parsimony and Bayesian inference. Based on the resulting phylogenies, a theory for the adaptive significance of the spore cyst and naked ascus within the bee habitat is proposed.

Methods

Morphological study

Spore cysts and asci were mounted in water on glass slides. Light photomicrographs were made on an Olympus AX70 Provis light microscope. Herbarium acronyms follow those of Index Herbariorum [36].

Collection

Ninety-six solitary bee nests were placed at eight localities on the island of Sjælland, Denmark. The nest holes ranged from 6 to 9 mm in diameter (to attract different species of cavity-nesting bees) and had a length of 19.5 cm.
The nests were placed 1–2.5 meters above the ground on the sides of buildings or below eaves with the entrances positioned to face southeast. Nests were opened and inspected annually from 2008 to 2012. Fungi growing on the pollen provisions, larvae, cocoons and nesting material were removed from a subset of the nests and identified. Overwintered honey bee (Apis mellifera) frames containing bee bread were collected in 2010 from three managed hives in Sjælland, Denmark. Bee bread with fungal growth resembling a spore cyst fungus was removed and the fungi isolated.

Isolation and cultivation
Isolates obtained for sequencing in this study were grown at 18 °C on a solid medium of malt agar with 20 % dextrose (MY20). Spore cysts of Bettsia alvei and the asci of Eremascus fertilis were plated directly onto MY20 and kept at 18 °C. Once sufficient growth was present to verify the identity of the fungus, a single hyphal tip was cut from each culture and transferred to a fresh plate. Specimens are deposited in the herbarium of The Natural History Museum of Denmark (C), and isolate cultures, given the designation “KVL ##-##” (Table 1) are the part of the permanent fungal isolate collection of the Insect Pathology laboratory in the Department of Plant and Environmental Sciences at the University of Copenhagen (formerly Den Kongelige Veterinær og Landbohøjskole, or KVL), where they are stored at −80 °C.

Molecular study
For two strains of Eremascus fertilis and two strains of Bettsia alvei (Table 1), genomic DNA was obtained by picking up ascocarps and mycelium from isolate cultures and grinding them inside a 1.5 ml Eppendorf tube. DNA was then isolated using the Qiagen DNeasy Plant Mini Kit (Hilden, Germany) using the standard protocol and eluted in two separate 50 µl fractions to avoid over-dilution.

The LSU and SSU regions were each amplified by PCR. Primers LR0R and LR7 [37] were used to amplify 1.4 kb of LSU, and primers NS1 and NS4 [38] were used to amplify 1.1 kb of SSU. PCR reactions were prepared in 50 µl volumes containing 29.8 µl of sterile deionized water, 5 µl of Taq polymerase reaction buffer (Sigma®), 1.0 µl 10 mM dNTP, 3.0 µl 25 mM MgCl₂, 0.2 µl Taq DNA polymerase (Sigma®), 5.0 µl each 10 µM primer and 1 µl of genomic DNA template. PCR was performed on a Biometra® thermocycler (Whatman) under the following conditions: for LSU: step 1) 1 min at 95 °C, 2) 1 min at 94 °C, 3) 30 sec at 51 °C, 4) 1 min at 72 °C, 5) return to step 2 34 times, 6) final step of 10 min at 72 °C; and for SSU: step 1) 1 min at 95 °C, 2) 1 min at 94 °C, 3) 30 sec at 51 °C, 4) 1 min at 72 °C, 5) 1 min at 94 °C, 6) 30 sec at 53 °C, 7) 65 sec at 72 °C, 8) return to step 2 34 times, 9) final step of 10 min at 72 °C. Samples were kept at 4 °C until electrophoresis was performed on 1 % agarose TAE gels and visualized with EZvision One® (Amresco). PCR products were cleaned using the Qiaquick® PCR purification kit (Qiagen) and were sent to Eurofins MWG Operon AG (Ebersberg, Germany) for sequencing. In addition to the amplification primers, LSU was sequenced with primers LR3R and LR5 [37], and SSU with primers NS2 [38] and SR7R (http://sites.biology.duke.edu/fungi/mycolab/primers.htm). Sequences were assembled using BioEdit [39].
For each region, a data matrix that included sequences from two isolates of *Eremascus fertilis*, two isolates of *Bettsia alvei*, and 63 other ascomycetes (Table 2) was assembled and manually aligned in MEGA5 [40]. Taxon sampling was focused on genera with reduced fruiting bodies. Many of the sequences came from James et al. [8] and were downloaded from the AFToL website (afterLog), while those from other studies [4, 41] came from GenBank. The LSU and SSU datasets were exported in NEXUS format and were combined in a single data file in PAUP* v. 4.0.10b [42]. The combined file was deposited in the Dryad Digital Repository, and can be accessed at http://dx.doi.org/10.5061/dryad.68s9o. This file was analyzed by maximum parsimony in PAUP*: using a random addition sequence and TBR swapping, 1000 heuristic replicates were performed, saving no more than ten best trees per replicate, followed by a final search of the saved trees. The file was also bootstrapped (2000 replicates) using the same search parameters except that only 10 heuristic replicates were performed per bootstrap replicate. A Bayesian analysis of the combined file was also performed using the program MrBayes v. 3.2 [43]. Based on the Akaike Information Criterion, the GTR + I + Γ model of DNA sequence evolution was selected as the best-fit model using the program Modeltest v. 3.06 [44]. A Markov chain Monte Carlo (MCMC) analysis was then run for 2,000,000 generations under the default settings, which was twice the number needed to keep the standard deviation of split frequencies below 0.01. Following Schoch et al. [4], Saccharomycotina was used as the outgroup for Pezizomycotina. Published single and multi-gene phylogenies [24, 45, 4] were followed in naming the major clades shown in Fig. 3.

**Results**

**Collection**

The predominant bee species in the solitary bee nests were *Osmia bicolor*, *O. leiana* and *Megachile centuncularis* and *M. versicolor*. Details of the contents of individual cells were recorded for 1553 of the approximately 8000 brood cells observed. Of 1553 brood cells, 1429 contained cocoons, 80 contained uneaten pollen provisions, and 182 had spore cysts on the pollen provisions, brood cell building materials, larval feces, larvae, or on the cocoon surface. Eighteen brood cells contained chalkbrood caused by *Ascosphaera*. Thirty-five of the 80 brood cells containing uneaten pollen provisions had *Bettsia* and *Ascosphaera* growing on and between the pollen grains. *Bettsia* and *Eremascus* were found growing on the bee bread of honey bee frames from two different localities (Table 1).

**Phylogenetic relationships of Eremascus and Bettsia**

**Molecular study**

The LSU matrix used for analysis included 862 characters, and the SSU matrix 1691 characters; the combined file thus included 2553 characters, of which 1762 were constant, 191 were variable but not parsimony-informative, and 600 were parsimony-informative. Two equally most-parsimonious trees of 3716 steps were recovered: a phylogram of one of these trees is shown in Fig. 3. Topologically, the parsimony and Bayesian trees agree with recent studies [21, 22]. Thus, Ascosphaeraceae as traditionally understood is also polyphyletic and the placement of *Ascosphaera* (BS = 87; posterior probability [PP] = 1) with *Byssosphaeria striatopora* (G.L. Barron & C. Booth) Arx and *Myxotrichum deflexum* Berk. *Eremascus albus*, the type species of the genus, was resolved in Eurotiomycetes, sister to *Ascosphaera* (BS = 82, PP = 1). *Bettsia* belonged to a fully supported clade including the leotiomycete genera *Pseudogyrospora*, *Leucoconiospora* and *Pseudoeurotiella*. Although the Leotiomycetes clade did not receive statistical support, the Eurotiomycetes clade was well supported (BS = 95, PP = 1).

*Eremascus* is a polyphyletic taxon. A new genus is needed for *E. fertilis* which is evidently related to *Myxotrichum* and *Byssosphaeria*. The placement of *E. fertilis* with Myxotrichaceae is supported morphologically by its narrow ellipsoid ascospores and uncoiled suspensors resembling stipitate asci (Fig. 2b-c). Narrow ascospores and stipitate asci are characteristic of Myxotrichaceae but are anomalous in the Onygenales (Eurotiomycetes), where *E. fertilis* was formerly placed [46].

The placement of *Bettsia alvei* in Leotiomycetes (Fig. 3) agrees with recent studies [21, 22]. Thus, Ascosphaeraceae as traditionally understood is also polyphyletic and the family must be circumscribed more narrowly to include *Ascosphaera* and *Arrhenospora*, but not *Bettsia*. *Bettsia* is most closely related to *Pseudoeurotiella*, *Pseudogyrospora* and *Leucoconiospora* in the family Pseudoeurotiellaceae. The cleistothecium of *Pseudoeurotiella* is formed by a
cellular peridium [47] rather than a double membrane as in a spore cyst, and the ascomal ontogeny of Pseudeurotium [48] is markedly different from that of a spore cyst [49, 50]. However, Pseudeurotium and Bettsia are similar in that they both have prototunicate asci and globose ascospores.

Table 2 Isolates and/or voucher specimens and GenBank accession numbers for LSU and SSU sequences

| Taxon                          | Isolate/strain/voucher specimen | GenBank ID/Sequence source |
|-------------------------------|---------------------------------|---------------------------|
|                               |                                 | LSU rDNA | SSU rDNA         |
| Acarospora schleicheri        | VR S-VII-98/30                  | AY640945 | AY640986         |
| Aleuria aurantia              | OSC 100018                      | AY544654 | NG_013139        |
| Anisomeridium polyomor        | 4237a                           | DQ782806 | DQ782877         |
| Ascosphaera apis              | CBS 402.96                      | FJ358275.1 | FJ358343        |
| Ascosphaera larvis            | ARSEF 7946                      | JX268535 | JX268535         |
| Aspergillus fumigatus         | ATCC 1022/JCM1738               | AY660917 | AB008401         |
| Aspergillus nidulans          | ATCC 16855/ FGSC4               | AF454167 | U77377           |
| Bettsia alvei                 | AA Wynns 5065 (C)               | KR139932 | KR139928         |
| Bettsia alvei                 | AA Wynns 5158 (C)               | KR139933 | KR139929         |
| Botryotinia fuckeliana        | OSC 100012                      | AY544651 | AY544695         |
| Byssosascus strictisporus     | CBS 642.66                      | AB040688 | AJ315170         |
| Bysschlamys nivea             | CBS 100.11                      | FJ358279 | FJ358345         |
| Caloscypha fulgens            | OSC 100062                      | DQ247799 | DQ247807         |
| Candida albicans              | SC5314                          | AACCQ0000290 | AACCQ0000290 |
| Candida tropicalis            | MUC130002                       | AAFN1000124 | M55527         |
| Capnodium coffeae             | CBS147.52                       | DQ247800 | DQ247808         |
| Capronia pilosella            | W. Untereiner WUC28             | DQ823099 | DQ823106         |
| Chlorociboria aeruginosa      | OSC 100056                      | AY544669 | AY544713         |
| Cladonia caroliniana          | F. M. Lutzoni 01-26-03.2        | AY584640 | AY584664         |
| Coccidiodium immitis          | ATCC 28868                      | AAEC0020 | AAEC0020         |
| Coccomyces dentatus           | OSC 100021                      | AY544657 | AY544701         |
| Cochliobolus heterostrophus   | CBS 134.39                      | AY544645 | AY544727         |
| Cudoniella clavus             | OSC 100054                      | DQ470944 | DQ470992         |
| Dactylella oxyspora           | CBS 280.70                      | AY902790 | AY902797         |
| Dermaea acerina               | CBS 161.38                      | DQ247801 | DQ247809         |
| Diaporthe eres                | CBS 109767                      | AF408350 | DQ471015         |
| Dothideomycotis sambuci       | DAOM 231303                     | AY544681 | AY544722         |
| Eleutherascus lactatii        | CBS 626.71                      | DQ168334 | DQ620997         |
| Endocarpon cf. pusillum       | S. Joneson 4028                 | DQ823097 | DQ823104         |
| Eremasculus albus             | UCB 50–026                      | M83258 | Q867787          |
| Eremascus albus               | CBS 975.69                      | FJ358283.1 | FJ358348.1     |
| Eremascus (Skoua) fertilis    | AA Wynns 5159 (C)               | KR139934 | KR139931         |
| Eremascus (Skoua) fertilis    | AA Wynns 5160 (C)               | HQ540515 | KR139930         |
| Exophiala pisciphila          | W. Untereiner WUC 137           | DQ823101 | DQ823108         |
| Geoglossum nigritum           | OSC 100009                      | AY544650 | AY544694         |
| Gymnoascus reessi             | CBS 259.61                      | FJ358284 | FJ358349         |
| Gyromitra californica         | OSC 100068                      | AY544673 | AY544717         |
| Histoplasma capsulatum        | ATFOL 1083                      | James et al. 2006 [8] | James et al. 2006 [8] |
| Hydropisphaera erubescens     | ATCC 36093                      | AYS45726 | AYS45722         |
Fig. 3 One of two equally most-parsimonious phylograms from a maximum parsimony analysis of nuclear ribosomal LSU and SSU DNA sequence data from 67 ascomycete fungi. Numbers above the branches are bootstrap support values (2000 replicates), and those below the branches are posterior probability values from a Bayesian analysis of the same data set. To the right of the arrows are diagrammatic illustrations of completely naked asci characteristic of *Eremascus albus*, *E. fertilis* (Pezizomycotina) and Saccharomycotina and nearly naked asci protected by hyphal wefts (*Gymnoascus*, *Myxotrichum*) or interspersed by sterile hyphae (*Eleutherascus*).
Taxonomy

**Skoua** A.A. Wynns, *gen. nov.*

Index Fungorum: IF551198

Type. *Skoua fertilis* (Stoppel) A.A. Wynns

Description Ascomata absent. Asci borne laterally from undifferentiated hyphae, prototunicate, subglobose, stipitate-like from two suspensors. Ascospores ellipsoidal, smooth, hyaline.

Etymology The generic name *Skoua* commemorates the Danish bee pathologist J. P. Skou, in acknowledgement of his major contribution to our understanding of spore cyst fungi.

**Skoua fertilis** (Stoppel) A.A. Wynns, *comb. nov.*

Index Fungorum: IF551199

Basionym. *Eremascus fertilis* Stoppel, Flora 97: 332. 1907.

Neotype of *Eremascus fertilis* (here designated): DENMARK: Zealand, Frederiksværk. Isolated from overwintering bee bread of *Apis mellifera* hive 74 belonging to Christian Pedersen, collected by A.A. Wynns #5159 (C). Index Fungorum: IF551197

Description Naked asci subglobose, 9–12 μm, on average 11 μm. Ascospores 4–8 × 3–5 μm. Natural habitat on bee bread inside the nests of *Apis mellifera*. In vitro growth at 18 °C, a low white mycelium on MY20, pale buff and radially sulcate with age (Fig. 2a), with abundant asci after four weeks. Conidia not observed.

Notes—In his description of *E. fertilis*, Stoppel [34] did not explicitly designate a holotype for this species; therefore, the ex-culture specimen A.A. Wynns #5159 deposited in herbarium C is here designated as the neotype.

Discussion

Convergent evolution of reduced fruiting bodies

Peridia, or the protective structures enclosing asci and ascospores, are diverse among the cleistothecial fungi. Peridia range from completely closed structures composed of many cells (e.g., *Pseudovestigium*), to cottony or cage-like enclosures of hyphae (e.g., *Byssocoschlamys, Myxotrichum, Gymnoascus*), to simple hyphae interspersed among naked asci and not forming an enclosure (e.g., *Byssosphaerus*). The morphology of the peridium was once thought to indicate relatedness [51] but hyphal or mesh-like peridia, cephalothecoid peridia, and now membranous peridia are known to have independently evolved in unrelated lineages and in taxa associated with insects [4, 5].

The results of the phylogenetic analyses (Fig. 3) unambiguously show that *Eremascus fertilis* is excluded from the class Eurotiomycetes and is not closely related to the type species of the genus, *E. albus*. *Eremascus fertilis* is therefore transferred to the new genus *Skoua*. This genus is closest to Myxotrichaceae, a family formerly placed in Eurotiomycetes but now considered to be a member of Leotiomycetales based on nrDNA-based phylogenies [45, 52]. Derived naked asci thus evolved at least twice within the Pezizomycotina: once in *Eremascus* (Eurotiomycetes) and once in *Skoua* (Leotiomycetes). Completely naked asci (i.e., without any vestigial peridial hyphae) occur elsewhere only in the basal lineages of Ascomycota (Taphrinomycotina and Saccharomyctina) [53]. Unlike the yeasts and yeast-like filamentous fungi (e.g., *Symbiotaphrina, Aureobasidium*), asexual reproduction by budding, fission, or otherwise does not occur in *Eremascus* and *Skoua*; therefore, the only yeast-like character of *Skoua* and *Eremascus* is naked asci.

Several taxa in Pezizomycotina (e.g., *Byssoschlamys, Gymnoascus, Pseudogymnoascus, Myxotrichum, Eleutherascus, Ascodesmis*) produce nearly naked asci that are enclosed by or interspersed with a loose network of hyphae (Fig. 3) known as a telaperidium or a reticuloperidium. Telaperidial and reticuloperidial ascomata are interpreted as loosely arranged cleistothecia or apothecia, a derived condition [46]. Interestingly, *Eremascus albus* and *Skoua fertilis* are allied to telaperidial and reticuloperidial taxa in both Eurotiomycetes (e.g., *Byssoschlamys, Gymnoascus*) and Leotiomycetidae (e.g., *Myxotrichum, Byssosphaerus*) (Fig. 3). More sequence data are needed to determine if the naked asci in *S. fertilis* evolved by reduction from a reticuloperidium or if reticuloperidial taxa evolved from *Skoua*-like ancestors. Reduced fruiting bodies are also found in *Ascodesmis* and *Eleutherascus* in Pezizomycotina, the most basal clade of the Pezizomycotina. In *Ascodesmis* the fruiting body consists of a bundle of naked asci.

Morphological convergence and dispersal

The discovery that the spore cyst and the naked asci have each evolved twice in unrelated lineages of bee-associated fungi suggests that these reduced fruiting bodies are well adapted to the bee habitat. Convergent evolution of fruiting body types and ascus types in unrelated fungi with similar dispersal strategies has occurred repeatedly in the Pezizomycotina, e.g., ascomata with reticuloperidial or cephalothecoid peridia [5, 54–56]. Evanescent asci or asci which break down to release the ascospores passively evolved multiple times in the Ascomycota and are correlated with insect dispersal [4, 57]. Similarly, the fragile peridium of the spore cyst fungi *Ascosphaera* and *Bettsia* and the evanescent naked asci of *Skoua* and *Eremascus* are interpreted as adaptive for dispersal by bees. Spore cysts break down when touched and dehisce with the activity of the bees, for example as bees move over and chew through the contents of brood cells in the process of emergence from their natal nests.
and during routine maintenance and construction of brood cells [58, 25, 59]. Disrupting the spore cyst membrane and subsequently picking up spores during emergence is a major contributor to the spread of chalkbrood in the managed solitary bee Megachile rotundata [60]. As the spore cysts are broken open, a sticky mucilage on the ascospores of Bettsia and Ascosphaera further ensures dispersal by the bee host [58, 25].

Spore cysts are not just broken open by the activity of bees, they also become impaled on their setae (Fig. 4). Impalement of fruiting bodies on arthropod setae is a spore dispersal mechanism found in other fungi, e.g. Myxotrichum (Leotiomycetes) and Auxarthron (Eurotiomycetes) [5]. In Myxotrichum and Auxarthron the mesh-like hyphal peridia function in attachment to the arthropod setae. Once impaled, ascospores fall through the holes in the mesh-like peridium and are dispersed as the insect (e.g., a fly) moves or flies around [5].

In this study, the peridia or membranous spore cyst walls of Ascosphaera and Bettsia growing within the nests of solitary bees remained intact in overwintering bee nests and in the absence of nesting activity. This is reminiscent of the spore dispersal strategy of Myxomycetes living in sheltered habitats. The peridium of Myxomycetes is dependent on the movement of invertebrates to disrupt the membrane, while in the absence of invertebrates the peridium remains intact for months [61].

The diversity and abundance of the spore cyst fungi is greater in the nests of solitary bees than in eusocial bees. The combination of the lack of social grooming and the nesting habits of solitary bees may contribute to the diversity and abundance of spore cyst fungi within the nests of these bees. In contrast to eusocial bees (e.g., honey bees), which may be active for months, solitary bees in temperate regions usually have only one generation per year and a relatively brief nesting and active period, sometimes as short as three weeks [62]. These nesting habits provide a stable environment for the slow-growing spore cyst fungi to establish and to develop mature ascocarps. On the other hand, because the active period of solitary bees is short compared with their eusocial relatives, the period for successful ascospore dispersal is much reduced. Ascospore release, coupled with the activity of the bees, maximizes spore dispersal of the fungus specifically by its host. Ensuring that ascospore release occurs in conjunction with the activity of the bees is particularly adaptive in the solitary bee habitat where the linear arrangement of brood cells means that siblings must pass through other brood cells to emerge from the nest (Fig. 1c–d).

Bee-mediated dispersal may be particularly critical for the spore cyst fungi and for Eremascus and Skoua because of their physiological requirements as xerophiles. Eremascus and Bettsia are included among the few xerophilic fungi called extreme xerophiles, meaning that they require rather than prefer a substrate with low water activity [63]. The bee habitat is a temporally static micro-environment that provides low water activity substrates, e.g., honey, bee bread, and pollen provisions, on which these slow growing xerophiles are able to thrive. A habitat with the combination of a temporally static environment and a provision of low water activity substrates is undoubtedly uncommon in nature and possibly unique to the nests of bees.

Conclusions
Microbial symbionts in bee nests and within the bees themselves may play a major role in maintaining bee health and in disease defense [64–67]. However, characterization of the microbial community in even the best-studied bee system, the honey bee, remains understudied, and its beneficial potential poorly understood [65]. Even less is known about the fungi intimately associated with the many species of solitary bees that make up the majority of the 20,000 bee species [68]; the literature is sparse [69] and the subject almost entirely unexplored. Understanding the phylogenetic
relationships of resident fungi within bee nests and their dispersal strategies may help to elucidate the role of these organisms in the bee habitat.

In this study, the systematic relationships of the bee specialist fungi are clarified and the monotypic genus *Skoua* is added to the microbial community associated with bees. The convergent evolution of spore cysts in *Bettsia* and *Ascosphaera*, and of naked asci in *Eremascus* and *Skoua*, is proposed to be adaptive for spore dispersal in the bee habitat. A taxonomic framework is provided for future studies of the understudied fungi occurring in this highly specialized niche.

Availability of supporting data
The data set supporting the results of this article is available in the Dryad Digital repository; doi:10.5061/dryad.6s80j, http://dx.doi.org/10.5061/dryad.6s80j.

Additional file

**Additional file 1: A supplementary figure in PDF format (pdf), 50 %-majority rule consensus phylogram from a Bayesian analysis of nuclear ribosomal LSU and SSU DNA sequence data from 67 ascomycete fungi.**

Competing interests
The author declares that they have no competing interests.

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