The Extent of Pollinator Sharing Among Fig Trees in Southern China

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Abstract

Background: The obligate mutualism between fig trees (Ficus, Moraceae) and pollinating fig wasps (Agaonidae) is a model system for studying co-evolution due to its perceived extreme specificity, but recent studies have reported a number of examples of trees pollinated by more than one fig wasp or sharing pollinators with other trees. This makes pollen flow between species and hybridization more likely. We reared pollinator fig wasps from figs of 13 Chinese fig tree species trees and established their identity using genetic methods in order to investigate the extent to which they are supporting more than one species of pollinator.

Results: Our results showed 1) pollinator sharing was frequent among closely-related dioecious species (where pollinator offspring and seeds develop on different trees), but not monoecious species and 2) that two pollinator species were developing in figs of one host tree species there was usually one fig wasp that was far rarer than the other. An exception was F. triloba, where its two pollinators were equally abundant.

Conclusions: Our results suggest that host expansion events where pollinators reproduce in figs other than those of their usual hosts are not uncommon among fig wasps associated with dioecious hosts. Because closely related trees typically have closely related pollinators that have a very similar appearance, the extent of pollinator-sharing has probably been underestimated. Any pollinators that enter female figs carrying heterospecific pollen could potentially generate hybrid seed, and the extent of hybridization and its significance may also have been underestimated.

Background

Ficus (Moraceae) is one of the most species-rich genera of woody plants in tropical and subtropical regions of the world (Harrison 2005), with more than 800 described species of free-standing trees, shrubs, climbers, and (hemi-)epiphytes (Corner 1965; Berg 1989; 1990). Figs are defined by their unique enclosed inflorescences (the fig or syconium) and their associated pollination system which requires entry into figs by highly-specialized fig wasps (Hymenoptera, Agaonidae). Pollinator fig wasps enter the figs to lay their eggs inside the ovules of the tiny flowers they contain. For a long period it was believed that each species of fig tree supported its own unique species of pollinator fig wasp, which was associated with no other Ficus species (Janzen 1979). When atypical pollinators were detected within figs it was assumed that these were rare mistakes that resulted in the death of the pollinators without the production of their offspring or generation of fertile seeds (Compton 1990; Ware and Compton 1992). More recently, it has been realized that more than one species of pollinator may be associated routinely with a single species of Ficus; and that widespread fig tree species can support multiple pollinators (Molbo et al. 2003; Haine et al. 2006; Sun et al. 2011; Chen et al. 2012; Yu et al. 2019). There are also examples of pollinator sharing, where two or more Ficus species are routinely hosts for a single species of fig wasp (Lopez-Vaamonde et al. 2002; Wachi et al. 2016; Wang et al. 2016). The one to one relationship that was originally envisaged is now realized to have been the result of the small number of host records available from each Ficus species, and their limited geographical coverage within the plants’ distributions, together with the close morphological similarities of closely-related pollinators making their identification difficult. Where two or more pollinators have been recorded as the routine pollinators of a single Ficus species they often appear to be associated with different habitats (Michaloud et al. 1996) and have allopatric or parapatric distributions within the ranges of their hosts. However, sampling intensity is again rarely sufficient to confirm this pattern of a single species of pollinator routinely servicing each Ficus species at any given location.

The assumption of extreme host specificity in fig wasps was based on a combination of the host records available and the apparent specialized co-adaptations required for a fig wasp to reproduce inside the figs of each Ficus species. Host choice by pollinators is made by the adult females and centres on long-distance plant-specific and developmental stage-specific volatile cues released by the figs when they are ready to be pollinated (van Noort et al. 1989; Grison-Pigé et al. 2002; Hossart-McKey et al. 2010). Pollinator females that arrive at a fig then need to be able to negotiate their way through a narrow ostiole in order to reach the flowers where they lay their eggs, and pollinator head shape is linked to the size of the ostiole (van Noort and Compton 1996). Successful oviposition once inside a fig depends on the fig wasp having an ovipositor that is longer than the styles through which its eggs are inserted (Nefdt and Compton 1996). Finally, successful development of their offspring depends on a galling response by the plant and gall forming insects are typically highly host specific (Weiblen 2004; Yu and Compton 2012; Ghana et al. 2015).

Although the relationship between fig trees and their pollinators is routinely described as a mutualism, the majority of Ficus species in Asia have a dioecious breeding system, where individual trees have figs that either produce only seeds (on ‘female’ trees) or only pollinator offspring (on ‘male’ trees) (Janzen 1979; Berg 2003). This situation contrasts with fig trees with a monoecious breeding system, where all the trees have figs that can produce both seeds and support the development of pollinator offspring. Monoecious fig trees are often large free-standing trees or stranglers (hemi-epiphytes) growing at low densities in forest habitats, whereas dioecious species are typically smaller and shrubby and more likely to have aggregated distributions (Berg 1990; Yang et al. 2015). Probably reflecting these differences, some pollinators of monoeocial species fly and transport pollen for long distances between trees (Ahmed et al. 2009), whereas the pollinators of dioecious Ficus species are believed to usually display more limited dispersal (Harrison and Rasplus 2006; Chen et al. 2011; Nazareno et al. 2013). The longer-distance dispersal exhibited by pollinators of monoeocial Ficus might be expected to increase the likelihood of two or more pollinator species developing in figs on a single tree, as has been reported in Africa (Compton et al. 2009), but host shifts are also likely to be easier between closely related species (Rasplus 1996) and there are numerous closely-related dioecious fig trees in Asia. Fig trees planted outside their normal range may also be more likely to support multiple pollinators, if their routine pollinators are absent locally (Corner 1965; Compton 1990; Patel et al. 1993).

The extent to which fig tree species growing within a single location are supporting more than one species of pollinator remains largely unknown, because most recent studies have concentrated on the pollination biology of individual species of fig trees (Chen et al. 2012; Darwell et al. 2014; Bain et al. 2016; Rodriguez et al. 2017; Yu et al. 2019). Pollinator sharing resulting in gene flow between closely-related Ficus species has nonetheless been detected (Wang et al. 2016). Here, we describe a Ficus community approach, where figs from southern China were screened for pollinator identity. The communities included mixtures of native and planted species and trees with both monoeocial and dioecious breeding systems. We address the following questions (1) is there any
difference on the extent of pollinator sharing between monoecious and dioecious Ficus? and (2) where two pollinators are present, does one species predominate?

Methods

Study site

Fig trees were sampled mainly at two sites in Guangdong province of SE China separated by about 200 kms: the South China Botanical Garden (N 23°10′46″, E113°21′06″; SCBG) with an area of 333 hectares and Dinghu Mountain (N23°09′21″, E112°30′39″; DHS) with an area of 1,133 hectares. Edaphic and climatic conditions at the two sites are similar and have a subtropical monsoon climate with distinct dry and wet seasons. The dry season runs from October to March, with 80% of annual precipitation concentrated in April-September. The mean annual temperature is 21.8 C in SCBG (Yu et al. 2006) and 21.9°C in DHS (Han et al. 2019), and the coldest mean monthly temperatures (13.1°C in SCBG and 12.6°C in DHS) occur in January.

More than 13,000 kinds of living plants are preserved in SCBG. The fig trees that support pollinators at SCBG include five monoecious figs, F. microcarpa, F. benjamina, F. subpisocarpa, F. virens, F. altissima, and seven dioecious fig trees, F. hirta, F. trifolba (one tree), F. auriculata, F. oligodon (two small trees), F. hispida, F. variegata var. chlorocarpa and F. pumila. In DHS, the natural vegetation comprises mainly southern subtropical monsoon evergreen broadleaved forests, reflecting moist local climatic conditions. The Ficus with pollinators present at DHS are F. microcarpa, F. benjamina, F. subpisocarpa, F. hirta, F. trifolba, F. hispida, F. fistulosa, F. variegata var. chlorocarpa, F. oligodon, F. erecta and F. pyriformis. The F. auriculata in SCBG are planted though the species is naturally distributed locally. The other dioecious Ficus at the two sites had not been planted, whereas the monoecious species had been planted.

The sampling area was extended so as to include three closely-related dioecious fig tree species belonging to subsection Frutescentiae of subgenus Ficus that have been recorded as sharing a single species of pollinator (Wiebes 1993). Pollinators were reared from figs of F. erecta var. beecheyana (Conghua E 113°57'9″; N 23°44'58″), F. pyriformis (DHS and Huizhou E 115°14′49″; N 23°5′49″ and Yangchun E111°47′9″; N22°10′23″). All were naturally-established individuals.

The identification of the pollinating wasps from each fig tree species

The pollinators of four monoecious and six dioecious fig tree species were identified using DNA sequencing (Table 1). Most non-pollinating fig wasp species (belonging to families other than Agaonidae) were excluded, but two species were utilized as out-groups. The four monoecious species are big trees with crops of more than ten thousand figs. F. microcarpa and F. benjamina belong to Urostigma subsection Conosycea, while F. subpisocarpa and F. virens belong to the subsection Urostigma. Among the dioecious species, F. hispida and F. fistulosa are small trees belonging to subgenus Sycomorus subsection Sycomorus (Cruaud et al. 2012). F. oligodon and F. auriculata are two closely related small trees belonging to subgenus Sycomorus, subsection Neomorpe (Berg 2004). F. hirta and F. trifolba belong subgenus Ficus, subsection Eriosycea. F. hirta is a shrub whereas F. trifolba is a small tree with larger crops. The three species belonging to subsection Frutescentiae are small shrubs. The monoecious species produce largely synchronous crops, whereas both sexes of the dioecious species usually exhibit asynchronous within-tree fruiting, with figs of different developmental stages present for longer periods on the plants.
The identities and haplotypes of pollinators reared from figs in Southern China.

| Ficus species         | Location | Insects genotyped | N trees | N figs | Pollinators                           | Haplotypes (frequency when >1) |
|-----------------------|----------|-------------------|---------|--------|--------------------------------------|-------------------------------|
| F. microcarpa         | SCBG     | 19                | 4       | 18     | Poll. 1. *Eupristina verticillata* agg. | H1(18), H2                     |
| F. benjamina          | SCBG     | 30                | 5       | 13     | Poll. 1. *Eupristina* sp. 1           | H1; H2(2); H3(13); H4(11); H5; H6; H7 |
| F. subpisocarpa       | SCBG     | 35                | 2       | 30     | Poll. 1. *Platyscapa cf. hsui* sp. 1  | H1(2); H2(2); H3; H4(2); H5; H6(2); H7(8); H8; H9; H10; H11; H12; H13; H14; H15(2); H16(2); H17(2); H18(2) |
|                       |          |                   |         |        | Poll. 2. *Platyscapa cf. hsui* sp. 2  | H1(2)                         |
| F. virens             | SCBG     | 8                 | 1       | 7      | Poll. 1. *Platyscapa coronata*        | H1(3); H2; H3(4)                |
| F. auriculata         | SCBG     | 31                | 1       | 13     | Poll. 1. *Ceratosolen cf. emarginatus* sp. 1 Poll. 2. *Ceratosolen cf. emarginatus* sp. 2 | H1; H2(29)                     |
|                       |          |                   |         |        |                                       | H1                             |
| F. oligodon           | DHS      | 29                | 1       | 8      | Poll. 1. *Ceratosolen cf. emarginatus* sp. 2 Poll. 2. *Ceratosolen cf. emarginatus* sp. 1 Poll. 3. *Blastophaga* sp 1 | H1(23); H2(2); H3               |
|                       |          |                   |         |        |                                       | H1                             |
|                       |          |                   |         |        |                                       | H1(2)                         |
| F. hispida            | SCBG     | 22                | 9       | 22     | Poll. 1. *Ceratosolen solmsi* marchali | H1; H2(20); H3                 |
| F. hispida            | DHS      | 6                 | 2       | 6      | Poll. 1. *Ceratosolen solmsi* marchali | H2(5); H4                     |
| F. fistulosa          | DHS      | 18                | 4       | 14     | Poll. 1. *Ceratosolen hewitti*        | H1(17); H2                    |
| F. hirta              | SCBG     | 8                 | 8       | 8      | Poll. 1. *Valisia javana hilli*        | H4(4); H6; H7; H8; H9           |
| F. hirta              | DHS      | 10                | 7       | 7      | Poll. 1. *Valisia javana hilli*        | H1(2); H2; H3(2); H4(4); H5     |
| F. triloba            | SCBG     | 1                 | 1       | 1      | Poll. 2. *Valisia javana hilli*        | H1                             |
| F. triloba            | DHS      | 21                | 9       | 17     | Poll. 1. *Valisia esquirolianae*       | H1(5); H2; H3(2); H4; H5        |
|                       |          |                   |         |        | Poll. 2. *Valisia javana hilli*        | H1(8); H2(3)                  |
| F. erecta var. beecheayana | Conghua | 29               | 6       | 25     | Poll. 1. *Blastophaga* sp 1            | H1(8); H2(14); H3; H4; H5; H6(3); H7 |
| F. pyriformis         | DHS      | 3                 | 1       | 3      | Poll. 1. *Blastophaga* sp 1            | H4, H9(2)                     |
| F. pyriformis         | Huizhou  | 6                 | 1       | 3      | Poll. 1. *Blastophaga* sp 1            | H1(2), H2, H4, H6(2)           |
| F. pyriformis         | Yangchun | 16                | 1       | 6      | Poll. 1. *Blastophaga* sp 1            | H2(4), H8(2), H9(9)            |
|                       |          |                   |         |        | Poll. 2. *Ceratosolen* sp 1            | H1                             |
| F. variolosa          | Dangan Island | 9               | 1       | 3      | Poll. 1. *Blastophaga* sp 1            | H1(9)                         |

**Fig wasp DNA extraction, amplification and analysis**

The mitochondrial genetic marker mtCOI was sequenced from an average of 23.2 fig wasp individuals reared from male figs of each *Ficus* species (range 8–35, total 301). Five fig wasp genera were represented (Table 1 and Table 2). All the sequenced fig wasps were adult offspring and therefore had developed successfully in the fig tree species from which they were reared.
Table 2

Pollinators are ranked by abundance from individual host tree species. The names of *Ficus* species are mainly from Flora of China v5 except that *F. triloba* is the revision of Berg 2007. The names of fig wasps are mainly based on Cruaud et al. 2012. *Eupristina verticillata* is now understood to be an aggregate of similar species. SCBG = South China Botanical Garden; DHS = Dinghu Mountain.

| Subsection | Species | Trees | Location | N figs | Pollinator sp. 1 | Pollinator sp. 2 | Pollinator sp. 3 |
|------------|---------|-------|----------|--------|-----------------|-----------------|-----------------|
|            |         |       |          |        | Identity N figs | Identity N figs | Identity N figs |
|            |         |       |          |        | (trees)         | (trees)         | (trees)         |
| Monoeicous | Conosycea | *F. microcarpa* | 3 | SCBG | 18 | *Eupristina verticillata* agg. | 18 | 19 (3) |
|            | F. benjamina | 5 | SCBG | 13 | *Eupristina* sp. 1 | 13 | 30 (5) |
| Urostigma  | *F. subpisocarpa* | 3 | SCBG | 21 | *Platyscapa* cf. *hsui* sp. 1 | 20 | 33 (3) |
|            | *F. virens* | 1 | SCBG | 7 | *Platyscapa coronata* | 7 | 8 (1) |
| Dioecious  | Neomorphe | *F. auriculata* | 1 | SCBG | 13 | *Ceratosolen* cf. *emarginatus* sp. 1 | 12 | 30 (1) |
|            |         | *F. oligodon* | 1 | DHS | 8 | *Ceratosolen* cf. *emarginatus* sp. 2 | 7 | 26(1) |
|            |         | *F. fistulosa* | 4 | DHS | 14 | *Ceratosolen* *hewitti* | 14 | 18 (4) |
|            | Eriosycea | *F. hirta* | 7 | DHS | 7 | *Valisia javana hilli* | 7 | 10 (7) |
|            |         | 8 | SCBG | 8 | *Valisia javana hilli* | 8 | 8 (8) |
|            | *F. triloba* | 9 | DHS | 17 | *Valisia javana hilli* | 8 | 11 (5) |
|            |         | 1 | SCBG | 1 | *Valisia javana hilli* | 1 | 1 (1) |
| Frutescentiae | *F. erecta* | 6 | Chonghua | 25 | *Blastophaga* sp. 1 | 25 | 29 (6) |
|            | *F. pyriformis* | 1 | Yangchun | 6 | *Blastophaga* sp. 1 | 5 | 15 (1) |
|            |         | 1 | Huizhou | 3 | *Blastophaga* sp. 1 | 3 | 6 (1) |
|            | *F. variolosa* | 1 | DHS | 1 | *Blastophaga* sp. 1 | 1 | 3 (1) |

Genomic DNA was extracted from the whole body of each fig wasp using the EasyPure Genomic DNA Extraction Kit (TransGen, Beijing, China). A 435–710 bp fragment of the mtCOI gene for each pollinating species was then sequenced following the protocol used in previous studies (Tian et al. 2015). The reaction was optimized and programmed on a MJ Thermal Cycler (PTC 200) as one cycle of denaturation at 94°C for 5 min, 35 cycles of 30 s denaturation at 94°C, 30 s at a 55°C annealing temperature, and 30 s extension at 72°C, followed by 8 min extension at 72°C. All amplified PCR products were purified using QIAquick spin columns (Qiagen) and were sequenced in an ABI 3730xl capillary sequencer using BigDye Terminator V 3.1 chemistry (Applied Biosystems). All unique haplotype sequences were deposited in GenBank (accession numbers: MW851213- MW851283).
We did not detect any indications of pseudo-genes, such as multiple peaks in chromatograms, stop codons or frame shift mutations (Song et al. 2008). Sequences were aligned using MUSCLE (Edgar 2004) implemented in MEGA 6.0 (Tamura et al. 2013) with manual corrections. DnaSP 5.0 was used to count the number of haplotypes (Librado and Rozas 2009).

Maximum likelihood trees were constructed using MEGA 6.0 (Tamura et al., 2013) for COI, and node supports were assessed based on 2000 bootstrap replicates. Kimura-2-parameter (K2P) distances within and between clades for COI haplotypes were then calculated. The clades with high gene sequence differences (larger than 0.02), were blasted to Genbank with the first 1–3 sequences sorted by percent identity. Two species of non-pollinating fig wasps reared from *F. hirta*, *Sycoscapter hirticola* (MG548706) and *Philotyposis josephi* (MG548673 and MG548674, both Pteromalidae) were included as outgroups (Yu et al. 2018).

**Results**

Phylogenetic analyses of the COI sequences detected 13 pollinator species that had reproduced within the figs of the 13 *Ficus* species, but there was not a 1:1 concordance between them. All the pollinator clades were strongly supported (Fig. 1; Table 1), with low within-clade and large between-clade K2P distances (Table S1) and the cumulative distribution of K2P distances indicating a marked barcoding gap between clades (Fig. 2). We therefore treat each clade as a distinct species. Based on the sequences downloaded from GenBank and our de novo sequencing we detected numerous examples of pollinators associated with more than one *Ficus* species and of *Ficus* species supporting the development of more than one species of pollinator. Up to three different species of pollinators were reared from the figs of a single host species and up to four host taxa were recorded for a single species of pollinator (Table 2).

The classical 1:1 pollinator and host *Ficus* relationship was only detected among two dioecious *Ficus* species (*F. hispida* and *F. fistulosa*), but it was the norm among the monoecious fig trees, where no pollinator-sharing was detected. *Ficus subpisocarpa* nonetheless supported the development of two closely-related fig wasps, rather than one (Table 2, Fig. 1). As reported previously based on molecular identifications, the same pollinator species (*Blastophaga* sp. 1) was reared from *F. erecta* var. *beecheyana*, *F. pyrifomis* and *F. variolosa*, but in addition the same species of fig wasp was also reared from figs of *F. oligodon*, an unrelated fig tree. *F. oligodon* was routinely supporting two species of *Ceratosolen*, both of which were shared with *F. auriculata* but no other species. The closely related taxa *F. hirta* and *F. triloba* also shared a pollinator (*Valisia javana hilli*) which was not reared from any other hosts.

Each fig wasps were generally reared from one or two host species (Table 2; Fig. 1). *Ficus* species supporting more than one species of fig wasp generally had one predominant pollinator that provided between 90–97% of the total reared individuals. The exception was *F. triloba* where its two pollinators were present in roughly equal proportions (Fig. 3; Table 1). Around half of the pollinators reared from *F. triloba* were *V. javana hilli*, a species routinely associated with *F. hirta* (*V. javana* complex sp. 1 in Yu et al. 2019).

**Discussion**

Our COI screening detected numerous examples of pollinator fig wasp species entering and successfully reproducing in more than a single host *Ficus* in southern China. Host overlap was frequent among pollinators of dioecious species and in most cases involved pairs of fig wasp species where one pollinator predominated and a second was reared only rarely. One interpretation of this is that the more rarely encountered pollinator species had other hosts where they were more abundant, but our screening across different *Ficus* species was not sufficiently extensive to confirm this and in some cases the pollinator species may simply be rare within our sampling area. Most examples of fig wasps developing in figs of more than one host involved fig trees that were closely related, but there were exceptions involving species of *Blastophaga* and *Ceratosolen* that were reared from figs normally associated with the other genus of pollinators. Fig wasp offspring developing successfully in unrelated host *Ficus* has been recorded previously from Africa (van Noort et al. 2013). This ability to develop inside hosts that are phylogenetically distant shows that the host specificity of fig wasps may be determined more by the choices made by searching adult females than by any physiological limitations.

Even within the *Ficus* species we sampled the size of the samples was not extensive and we are unlikely to have detected the full range of *Ficus* hosts being utilized locally by the fig wasps. There was generally one routine pollinator species combined with rarer entries by two or more additional pollinators (Moe et al. 2011; Yang et al. 2015). An exception to the general pattern of pollinator sharing where one pollinator species predominated was provided by *F. triloba*, where two pollinator species were present in roughly equal numbers of its figs, but more samples, taken throughout the year, will be needed to confirm this pattern. Some of the trees we sampled were planted individuals and this may have increased the extent of pollinator sharing that we detected. Our results nonetheless suggest that exceptions to the ‘classical’ one pollinator to one tree relationship are routine among sympatric dioecious fig tree species in southern China, to the extent that among trees with this breeding system strict specifcity is the exception, not the norm.

Fig wasps develop inside figs on male trees of dioecious fig species, but it is likely that similar entry by two or more pollinator species is taking place in both male and female figs. Pollinator host choice, based mainly around species-specific volatile attractants released by receptive figs, is the major isolating mechanism that helps prevent heterospecific pollen being deposited on the flowers inside female figs, but is not always effective (Souto-Vilarós et al. 2018). Other isolating mechanisms such as pollen incompatibility appear to be poorly developed in *Ficus* (Huang et al. 2019), so whenever fig wasps species are entering female figs of two more host trees in an area there is the possibility of viable hybrid seed being developed. The extent to which hybrids can mature successfully and can lead to introgression between species is unknown, but mature natural hybrid individuals are known to occur (Wilde et al. 2020). Some artificially generated hybrids appear to be at no reproductive disadvantage in terms of seed production, but male hybrid offspring can be sterile because pollinators cannot develop inside their figs (Ghana et al. 2015, but see also Yakushijii 2012), so patterns of introgression may be complex.

Sharing of pollinators was not a feature of the monoecious fig tree species we sampled. This is not the case elsewhere, where for example up to four different species of pollinators from figs of a single individual tree in Africa (Compton et al. 2009). Monoecious and dioecious fig trees differ in numerous ways that may influence pollinator behavior including growth form (trees versus shrubs), flowering phenology (large synchronous crops versus smaller asynchronous
crops) and typical local abundance (monoecious species are often more dispersed). Perhaps more significantly in our study area and across SE Asia, there an exceptionally high diversity of dioecious species, most of which are pollinated by fig wasps that belong to a small number of genera. Opportunities for chance landing on figs of atypical hosts are therefore greater for those insects associated with dioecious hosts, but in addition most of the sharing of pollinators was between closely related dioecious species, which are likely to be generating relatively similar attractant cues (Wei et al. 2014; Wang et al. 2016). The morphology of closely-related pollinator fig wasps is often very similar, and our results emphasize that pollinator-sharing is likely to have been underestimated because of this. Barcoding and other molecular identification techniques are used increasingly to distinguish between fig wasp species, but our results also highlight the need to sequence fig wasps from several figs, even if they look alike, in order to detect pollinator species that may be present at low frequencies. More than one morphologically similar species can even be reproducing within the same individual figs (Sutton et al. 2017).

Conclusions

Our survey of the fig wasp pollinators associated with local assemblages of *Ficus* species in Southern China revealed contrasting pollinator relationships between monoecious and dioecious trees. Monoecious trees and their pollinators largely displayed a highly specific one pollinator for one tree association. Among dioecious species there was no such specificity, with frequent sharing of pollinators across trees and two or more species of pollinators associated with each tree species. Possible biological traits favoring this breakdown in pollinator specificity among dioecious *Ficus* include their extended asynchronous flowering phenologies and the mixtures of closely-related species that can grow in close proximity. This lack of specificity suggests that the extent of pollen flow between dioecious fig tree species is likely to have been underestimated, with unknown consequences.

Abbreviations

SCBG
South China Botanical Garden
DHS
Dinghu Mountain
DNA
Deoxyribonucleic acid
mtCOI
Mitochondrial cytochrome oxidase I
COI
Cytochrome oxidase I
K2P
Kimura-2-parameter

Declarations

Author's contributions

YH designed the research and analyzed data. LYL, CYF and JYX performed laboratory work. CYF also collected samples. KFS analyzed data. YH, CS and SGC co-wrote the manuscript.

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Availability of data and materials

All unique haplotype sequences of COI sequence for all the sampled pollinators were deposited in GenBank (accession numbers: MW851213 - MW851283)

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no conflict interests.

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Figures

Figure 1

COI ML phylogenetic tree of the pollinators of the sympatric figs, with sequences of two non pollinators (Pteromalidae) as outgroups. Node support rates are shown. Haplotypes are also listed together with their host figs.
Figure 2
Cumulative distribution of Kimura pairwise genetic distances (K2P) for COI of pollinators associated with Ficus species. Intraspecific distance ranged between 0-1.3% and interspecific distances ranged from 3.3-29.2%. A marked barcoding gap separated the within- and between-putative species pairwise distances.
Figure 3

The proportion of different pollinator species in each Ficus species. The columns with red marks show the same major pollinator species in F. hirta and F. triloba; while those with blue marks show the same major pollinator species of F. erecta var. beecheyana, F. pyriformis and F. variolos.

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