Remote tropical island colonization does not preclude symbiotic specialists: new evidence of mycorrhizal specificity across the geographic distribution of the Hawaiian endemic orchid Anoectochilus sandvicensis

Sean Swift1, Sherilyn Munroe1, Chaewon Im2, Laura Tipton1 and Nicole A. Hynson3*

1Department of Botany, University of Hawaii Manoa, 3190 Maile Way, Honolulu, HI 96822 USA, 2John A. Burns School of Medicine, University of Hawaii Manoa, 651 Ilalo street, Honolulu, HI 96813 USA and 3Pacific Biosciences Research Center, University of Hawaii Manoa, 3190 Maile Way, Honolulu, HI 96822 USA

*For correspondence. E-mail nhynson@hawaii.edu

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INTRODUCTION

Carlquist in his classic work Island biology (1974) posited that the successful colonization of islands by obligate symbiotic organisms such as insect-pollinated plants, would favour generalists. Recent studies of plant–pollinator networks on islands provide some empirical support for Carlquist’s hypothesis, where native and endemic plants along with their pollinators tend to be generalists and less specific than expected by chance (Castro-Urgal and Traveset, 2014; Trojelsgaard et al., 2015; Hervias-Parejo and Traveset, 2018). However, the degree of symbiotic specificity across the geographic distributions of island native, and especially endemic, biotic organisms such as insect-pollinated plants, would favour generalists. Plants that form obligate symbiotic associations with microbes dominate island ecosystems, but the relationship between island inhabitance and symbiotic specificity is unclear, especially in the tropics. To fill this gap, we examined the mycorrhizal specificity of the Hawaiian endemic orchid Anoectochilus sandvicensis across multiple populations encompassing its entire geographic distribution.

METHODS

By molecular phylogenetic approaches we identified the mycorrhizal fungi associated with A. sandvicensis across its entire geographic distribution and determined the relationship of these fungi to others found elsewhere around the globe. With richness estimators, we assessed the mycorrhizal specificity of A. sandvicensis within and among islands. We then tested whether geographic proximity of orchid populations was a significant predictor for the presence of particular mycorrhizal fungi and their community composition.

RESULTS

We found that each population of A. sandvicensis forms specific associations with one of three fungi in the genus Ceratobasidium and that the closest relatives of these fungi are globally widespread. Based on diversity indices, A. sandvicensis populations were estimated to partner with one to four mycorrhizal taxa with an estimated total of four compatible mycorrhizal fungi across its entire distribution. However, the geographic proximity of orchid populations was not a significant predictor of mycorrhizal fungal community composition.

CONCLUSIONS

Our findings indicate that the colonization and survival of plant species on even the most remote oceanic islands is not restricted to symbiotic generalists, and that partnering with few, but cosmopolitan microbial symbionts is an alternative means for successful island establishment. We suggest that the spatial distribution and abundance of symbionts in addition to island age, size and isolation should also be taken into consideration for predictions of island biodiversity.

Key words: Anoectochilus sandvicensis, Ceratobasidium, island biogeography, mycorrhizal fungi, Orchidaceae, rhizoctonia, tropical ecology.
as the rhizoctonias, which includes species in the families Tulasnellaceae, Ceratobasidiaceae and Serendipitaceae (Dearnaley et al., 2016; Weiβ et al., 2016). Unlike other lineages of mycorrhizal fungi, some orchid-associated species are thought to be capable of living independently of a host as soil saprotrophs (Veldre et al., 2013; Weiβ et al., 2016), indicating that their geographic distributions may not be strictly tied to those of their hosts (McCormick and Jacquemyn, 2014). Based on a recent biogeographic report, orchid-associated Ceratobasidiaceae and Tulasnellaceae appear to be globally widespread, while Serendipitaceae species are less common (Jacquemyn et al., 2017). However, distribution data on orchid mycorrhizal fungi are patchy, especially for the tropics where some of the greatest orchid diversity occurs (Swarts and Dixon, 2009).

Symbiotic diversity in orchids can range from highly specific associations with single species of fungi, to much more general interactions with multiple fungal lineages (Shefferson et al., 2005; McCormick et al., 2006; Dearnaley, 2007; Waterman and Bidartondo, 2008; Jacquemyn et al., 2010). While information on the mycorrhizal associations of island-inhabiting orchids is limited, previous studies provide mixed support for Carlquist’s predictions. Some studies have found high degrees of mycorrhizal specificity among native orchids, while others have found them to be generalists associating with a diversity of rhizoctonias, but favouring Tulasnellaceae species (Otero et al., 2002; Ma et al., 2003; Kristiansen et al., 2004; Otero et al., 2004; Martos et al., 2012; Jacquemyn et al., 2017). Information is lacking particularly on the mycorrhizal specificity and associations of tropical island orchids, many of which are threatened or endangered by extinction (Swarts and Dixon, 2009). Furthermore, previous studies are limited in their ability to examine explicitly the relationship between island inhabitance by orchids and mycorrhizal specificity due to: (1) many of the orchid species examined thus far have distributions that include islands and continents; and (2) when mycorrhizal specificity has been detected in island endemic orchids, these observations are limited to one or two orchid populations. Here we leverage the presence of an island orchid Anoeoctochilus sandvicensis which occupies six discrete oceanic islands to examine the degree of symbiotic specificity among multiple host populations and across the host’s entire geographic distribution.

The Hawaiian Islands provide a unique backdrop for biogeographic studies. Hawai‘i is the most isolated archipelago on the planet, made up of eight main islands that range in age and size from 0.4 to 5.1 million years old and from 115.5 to 10,432 km², respectively (Ziegler, 2002). From Ni‘ihau, the oldest and furthest west of the main Hawaiian Islands, to the youngest and furthest east Island of Hawai‘i spans approx. 620 km, while only 11 km separate the closest main islands of Maui and Kaho‘olawe; each island also harbours unique biomes ranging from tropical rain forest to volcanic deserts. Accordingly, the species that have arrived there have done so by improbable means and many have undergone rapid radiations (Ziegler, 2002). This has resulted in unique flora and fauna, with an estimated 89% of Hawai‘i’s vascular plant species found nowhere else in the world (Wagner et al., 1999). Despite their propensity for long-distance dispersal (Givnish et al., 2016), Hawai‘i harbours only three distantly related native orchids, including A. sandvicensis (tribe Cranichideae) (Wagner et al., 1999).

Due to Hawai‘i’s extreme isolation and the requirement of all orchids for symbiotic mycorrhizal fungi, we predicted that the orchids that inhabit these islands should be mycorrhizal generalists (sensu Carlquist, 1974). Specifically, we set out to test the hypotheses that: (1) A. sandvicensis partners with a diversity of mycorrhizal fungal lineages across its geographic distribution; and (2) as distance among A. sandvicensis populations increases, the similarity of mycorrhizal fungal communities decreases.

The fact that A. sandvicensis is endemic and only found on the main Hawaiian Islands makes it an interesting study system to examine the interactions between endemism and symbiont specificity. Due to the isolation of Hawai‘i, one might assume that endemic species that require mutualists may, by default, be specialists due to a limited number of available and compatible symbionts. However, this appears not to be the case for island endemic plants and their pollinators (Castro-Urgal and Traveset, 2014; Trojelsgaard et al., 2015; Hervias-Parejo et al., 2018). While there is conflicting information on the degree of mycorrhizal specificity among endemic vs. more widespread orchids (Swarts et al., 2010; Phillips et al., 2011; Jacquemyn et al., 2016) based on evidence from relatives of A. sandvicensis, there is some support that this genus is capable of forming mycorrhizae with a diversity of rhizoctonia fungi, and even one normally saprotrophic species of Mycena (Guo et al., 1997; Jiang et al., 2015). Of the three endemic Hawaiian orchids, we chose to focus on A. sandvicensis because, while locally rare, it is the most common native orchid species in Hawai‘i found throughout all the major islands including Maui, O‘ahu, Hawai‘i, Kaua‘i, Moloka‘i and Lāna‘i, and its habitats range from montane wet tropical forests to mesic shrublands. We used a molecular and phylogenetic approach to determine the identities and evolutionary relatedness of the mycorrhizal fungi associated with 12 populations of A. sandvicensis from four islands representing the full extent of its geographic distribution. Using richness estimators, we assessed the mycorrhizal specificity of A. sandvicensis among populations and across its distribution, and then tested the relationship between geographic distance among orchid populations and mycorrhizal fungal community similarity.

**MATERIALS AND METHODS**

**Field sites and sampling**

With the assistance of local botanists, populations or clusters of Anoeoctochilus sandvicensis were located on the islands of O‘ahu, Maui, Kaua‘i and Hawai‘i (Table 1). Once research permits were obtained for each site, between July 2013 and April 2017 we sampled roots from 12 A. sandvicensis populations (AN1–AN12) across the four islands (three on O‘ahu, four on Maui, three or four on Kaua‘i and one on Hawai‘i Island, Table 1). In general, populations were separated by >100 m, except for those found on the steep and difficult to access Kahili Ridge, Kaua‘i where the two closest (AN8 and AN10) were 1.8 m apart, making it challenging to confirm in situ that they were separate populations. Measured using the GPS...
Table 1. Orchid population site locations for samples collected in this study and fungal operational taxonomic units (OTUs) identified from each population including new accession numbers and best BLAST match from NCBI’s GenBank, and number of representative sequences per OTU

| Location                  | Population | Elevation (m) | Fungi identified          | Accession number | Best BLAST match name and accession number                                      | No. of representative sequences | Maximum score | Query coverage (%) | Per cent identity (%) |
|---------------------------|------------|---------------|---------------------------|------------------|--------------------------------------------------------------------------------|--------------------------------|---------------|---------------------|------------------------|
| Mt. Ka’ala, O’ahu         | AN01       | 1208          | Ceratobasidium sp. 1      | MF539823         | Ceratobasidium sp. AB290022.1                                                | 21                             | 1046          | 100                 | 98                     |
|                           |            |               | Tulasiellaceae sp.        |                  | Tulasiella sp. JN65638.1                                                      | 2                              | 1119          | 98                  | 90                     |
|                           |            |               | Polyporales sp. 1         |                  | Polyporales tricholoma KP943502.1                                            | 1                              | 398           | 83                  | 81                     |
|                           |            |               | Polyporaceae sp.          |                  | Coriolopsis rigida JF894112.1                                                 | 1                              | 1079          | 93                  | 99                     |
|                           | AN02       | 1195          | Ceratobasidium sp. 2      | MF539822         | Ceratobasidium sp. AB290022.1                                                | 9                              | 1138          | 100                 | 98                     |
|                           | AN03       | 1227          | Ceratobasidium sp. 2      | MF539822         | Ceratobasidium sp. JX138539.1                                                 | 19                             | 1138          | 100                 | 98                     |
|                           |            |               | Polyporales sp.           |                  | Polyporales tricholoma KP943502.1                                            | 2                              | 398           | 83                  | 81                     |
| Waikamoi Nature Preserve, Maui | AN04      | 1392          | Ceratobasidium sp. 1      | MF539823         | Ceratobasidium sp. AB290022.1                                                | 5                              | 1046          | 100                 | 98                     |
|                           |            |               | Trechisporales sp.        |                  | Subulicystidium perlongisporum KP268489.1                                     | 4                              | 327           | 99                  | 85                     |
|                           | AN05       | 1447          | Galerina sp.              | MF539823         | Galerina sp. JX029941.1                                                        | 1                              | 1050          | 100                 | 94                     |
|                           |            |               | Ceratobasidium sp. 1      |                  | Ceratobasidium sp. AB290022.1                                                | 30                             | 1046          | 100                 | 98                     |
|                           | AN06       | 1476          | Galerina sp.              | MF539823         | Galerina sp. JX029941.1                                                        | 1                              | 1046          | 100                 | 98                     |
|                           |            |               | Ceratobasidium sp. 1      |                  | Ceratobasidium sp. AB290022.1                                                | 26                             | 1046          | 100                 | 98                     |
|                           | AN07       | 1391          | Ceratobasidium sp. 1      | MF539823         | Ceratobasidium sp. AB290022.1                                                | 21                             | 1046          | 100                 | 98                     |
| Kahili Ridge, Kaua’i      | AN08       | 829           | Ceratobasidium sp. 1      | MF539823         | Ceratobasidium sp. AB290022.1                                                | 7                              | 1046          | 100                 | 98                     |
|                           | AN09       | 810           | Ceratobasidium sp. 1      | MF539823         | Ceratobasidium sp. AB290022.1                                                | 12                             | 1046          | 100                 | 98                     |
|                           |            |               | Athelieaeae sp.           |                  | Athelieaeae sp. JX898945.1                                                     | 3                              | 1046          | 100                 | 98                     |
|                           | AN10       | 830           | Ceratobasidium sp. 1      | MF539823         | Ceratobasidium sp. AB290022.1                                                | 19                             | 1046          | 100                 | 98                     |
| Wainiha Valley, Kaua’i    | AN11       | 823           | Ceratobasidium sp. 3      | MF539824         | Rhizoctonia sp. JQ85895.1                                                     | 18                             | 924           | 100                 | 96                     |
| Ola’a tract, Volcano National Park, Hawai’i | AN12      | 1162          | Ceratobasidium sp. 2      | MF539822         | Ceratobasidium sp. JX138539.1                                                 | 35                             | 1138          | 100                 | 98                     |
coordinates of our sampling sites, the greatest distance between populations was approx. 532 km between AN11 from Wainiha Valley on Kaua‘i and AN12 from the Ola‘a tract in Volcanos National Park on Hawai‘i Island (Table 1). Vegetation types for orchid collection sites were lowland to montane, wet to mesic, native forests and shrublands.

Using a trowel cleaned with 70 % ethanol, we excavated small rhizome and root sections from each population of A. sandvicensis. Due to the rhizomatous and mat-forming growth habit of A. sandvicensis, we were unable to determine individuals in the field, so root samples are representative of clusters of intertwined individuals no more than 0.5 m apart. Root samples between 3 and 6 cm long were placed into plastic bags and put directly on ice. After returning from the field, samples were washed with tap water and sectioned with clean razor blades into small sub-samples approx. 1–2 cm long that were stored in 600 μL of cetyltrimethylammonium bromide (CTAB) buffer. These samples were frozen at 0 °C until they were returned to the University of Hawai‘i at Mānoa where they were stored at –20 °C for future molecular analyses.

**Root sectioning**

Replicate root samples of A. sandvicensis from each population were removed from CTAB buffer, surface sterilized in a 10 % bleach solution for 30 s and suspended in water immediately prior to sectioning. Each root sample was sub-divided into 0.5 mm transverse sections. For each root sub-sample, 3–4 sections that showed signs of colonization (dense fungal pelotons in cortical cells, Fig. 1) were selected for DNA isolation. Colonized sections from each root sub-sample were placed in 2 mL bead beating tubes with 1000 μL of CTAB and sterile 2 mm and 5 mm glass beads, and stored at –20 °C prior to DNA isolation. We extracted DNA from a minimum of 16 samples per orchid population.

**Molecular methods**

Colonized root sections suspended in CTAB buffer were thawed at 65 °C and frozen in liquid nitrogen three times to soften tissue prior to bead homogenization. Samples were homogenized in a mini beadbeater (Biospec) and then incubated at 65 °C for 1 h and mixed by inverting every 20 min. DNA was extracted using the Qiagen DNeasy Blood & Tissue kit (Qiagen Inc., Valencia, CA, USA) following a modified protocol (Hyinson and Bruns, 2009). Using PCR, the nuclear ribosomal internal transcribed spacer (ITS) region was amplified with the orchid mycorrhizae optimized primer combination ITS1OF/ITS4OF and, for a sub-set of samples, the primer combination ITS1/ITS4-TUL and the PCR conditions described in Taylor and McCormick (2008). For representatives of each Ceratobasidium operational taxonomic unit (OTU; see the Results), a larger fragment of the ribosomal large sub-unit (LSU) was amplified using the primer pair ITS1OF (Taylor and McCormick, 2008) and TW14 (Bidartondo and Duckett, 2009). PCR products were cleaned using ExoSap-IT (Affymetrix). Clean PCR products were unidirectionally sequenced using the primer ITS1OF for ITS amplicons and LROR for LSU amplicons. Sanger sequencing was performed by either Genewiz (Cambridge, MA, USA) or ASGPB (University of Hawai‘i, Honolulu, HI, USA).

**Fungal identification**

Sequences were trimmed in Geneious (v.5.4.6). OTUs were binned at 97 % identity using the QIIME ‘open reference’ workflow (v1.9.1; Caporaso et al., 2010). Conserved regions flanking ITS1 were removed using ITSx (v1.0.11; Bengtsson-Palme et al., 2013). Open reference clustering was performed against the dynamic UNITE fungal database version 28.06.2017. Sumaclust was used during open reference OTU binning (Mercier et al., 2013), and SortMeRNA (Kopylova et al., 2016) was used for subsequent de novo OTU binning. OTU designation for sequences that contained ambiguous sites was confirmed by local alignment using MUSCLE (v. 3.8.31). Initial taxonomic assignments were based on Blast alignments with the non-redundant databases of NCBI and UNITE (Nilsson et al., 2015). Queried sequences were considered a match at the generic level if the query coverage was 100 % and percent identity was ≥94 %, and a match at the familial or ordinal levels if the query coverage was ≥93 % or ≥83 %, and if percent identity was ≥90 % or ≥81 %, respectively (Table 1). Representative samples with ITS sequences matching fungi from known orchid mycorrhizal groups, specifically the family Ceratobasidiaceae or the genus Ceratobasidium, were resequenced with the LSU primer LROR for phylogenetic reconstruction. Representative ITS sequences for all OTUs and representative LSU sequences for Ceratobasidiaceae OTUs were deposited in GenBank (MF539822:MF539824; MF599203:MF599205 and MF574046:MF574052).

**Phylogenetic analyses**

For phylogenetic reconstruction, ITS and LSU sequences were selected for each Ceratobasidium or Ceratobasidiaceae OTU based on sequence length and quality. Phylogenetic analyses were performed on two separate data sets. The first data set was comprised of concatenated ITS/LSU sequences representing each OTU and reference sequences of Rhizoctonia anamorphs in Ceratobasidiaceae from recently published phylogenies (Gonzalez et al., 2001, 2016). Sequences of Hydnum and Clavulinia were included as an outgroup. The second data set
Estimates of mycorrhizal specificity and distance decay relationship of mycorrhizal fungi among orchid populations

To assess whether our sampling efforts resulted in detection of the full range of fungi that associate with A. sandvicensis, we calculated the first three Hill numbers using the iNext package v 2.0.12 in R (Chao et al., 2014). Because values of observed species richness often underestimate total richness (Hughes et al., 2001), Hill numbers offer numerous advantages over reporting observed species richness alone as well as advantages over other richness estimators (Chao et al., 2014). The first Hill number ($q = 0$) is simply an estimate of richness without regard for relative abundance. The second two Hill numbers ($q = 1$ and $q = 2$) are the exponential of Shannon entropy and the inverse Simpson concentration, respectively, which both incorporate species (or OTU) abundance into their estimates, but Shannon is the effective number of common OTUs whereas Simpson is the effective number of dominant OTUs in the assemblage. We used the number of sequences of each mycorrhizal OTU as a measure of taxon abundance. We estimated mycorrhizal richness and diversity both by A. sandvicensis population (Fig. 2A) and by island (Fig. 2B).

To test whether geographic proximity of A. sandvicensis populations was a significant predictor of mycorrhizal fungal community composition or the presence of particular Ceratobasidium OTUs, we performed Mantel correlation tests between geographic distance (in metres) and Bray–Curtis community dissimilarity for either all mycorrhizal taxa or only Ceratobasidium OTUs. Mantel test $P$-values were calculated based on 1000 permutations in the R package vegan version 2.4–6.

RESULTS

The dominant fungal genus associated with populations of Anoectochilus sandvicensis was Ceratobasidium, where each orchid population partnered with a single OTU in this genus (Table 1). Overall, Ceratobasidium sequences accounted for 94% of the total 237 sequences from this study, and in seven out of the 12 orchid populations sampled it was the only fungus detected. The additional fungi found in the remaining five populations included saprotrophs (two polypore OTUs represented by one and three sequences, respectively; two sequences of a single Galerina species, and four sequences of a single OTU in the order Trechisporales, Table 1). We also detected two additional putative mycorrhizal OTUs including the same Atheliaceae taxon (100% query coverage and 99% identity) found associating with non-native pines in plantations on Maui (Hynson et al., 2013), here represented by three sequences, and a single Tulasnellaceae OTU represented by two sequences.
(Table 1). This Tulasnellaceae species was amplified with the ITS1OF/ITS4OF primers. None of our samples amplified fungi with the ITS1/ITS4-TUL primers. Overall, we had a PCR success rate of 87.9% with the ITS1OF/ITS4OF primers.

Based on our binning of sequences at 97% similarity, we detected three Ceratobasidium OTUs (Table 1). By far the most common was Ceratobasidium sp. 1, which was detected in eight populations of A. sandvicensis on three islands and represented by a total of 141 sequences (Table 1). Ceratobasidium sp. 2 was the next most common, found associating with three populations across two islands (Table 1) represented by 63 sequences, while Ceratobasidium sp. 3 was only detected from a single orchid population in the Wainiha Valley, Kaua‘i and represented by 18 sequences. Each of the 12 orchid populations partnered with a single OTU of Ceratobasidium, and all sites except Mt. Ka‘ala on O‘ahu harboured only a single Ceratobasidium OTU (Table 1).

The concatenated ITS + LSU phylogeny (Fig. 3) included expert curated reference sequences from Ceratobasidium anamorph group AG and indicated that all three Ceratobasidium OTUs from our study formed a monophyletic clade with two Ceratobasidium OTUs in the AG-H and AG-R anamorph groups from Japan and the USA, and another Ceratobasidium OTU (isolate CBS 148.54) from France (Fig. 3). Along with representative sequences from each of the three OTUs described in the current study, the concatenated data matrix included an additional 30 taxa from recent phylogenies of Rhizoctonia anamorphs within Ceratobasidiaceae (González et al., 2001, 2016). The final alignment contained 1755 characters of which 978 were included in the analysis. Of the included characters, 239 were parsimony informative.

The ITS phylogeny provided a finer scale phylogenetic context and was annotated with the geographic origin and mycorrhizal status of each sequence (Fig. 4). This data set included representative sequences from each of our three Ceratobasidium OTUs and 69 sequences of Ceratobasidium taxa from GenBank. The final alignment contained 1664 characters of which 533 were included in the analysis. Of the included characters, 105 were parsimony informative. The maximum pairwise distance among ITS sequences included in the analysis was 8.33%. Based on our ITS phylogeny, the three Ceratobasidium OTUs described from A. sandvicensis did not form a monophyletic group (Fig. 4). Ceratobasidium sp. 2 formed a well-supported sub-clade with sequences of Ceratobasidium from a previous study of orchid mycorrhizal fungi in Japan (bootstrap support = 96; Shefferson et al., 2010). Ceratobasidium spp. 1 and 3 were retained in the in-group but did not form well-supported sub-clades with each other or with any of the reference sequences included in the analysis (Fig. 3). The best supported clade that includes Ceratobasidium spp. 1 and 3 also includes Ceratobasidium sp. 2 as well as a suite of other taxa in Ceratobasidiaceae including both orchid mycorrhizal and non-mycorrhizal species, and were found in Argentina, Australia, the Azores archipelago, China, France, Italy, Japan, South Africa, Spain and the USA (Fig. 4).

Based on our calculations of the first three Hill numbers, we estimated that the total mycorrhizal fungal richness ($q = 0$) for all A. sandvicensis populations was 3.92 ± 1.96 (s.e.), whereas exponential Shannon’s diversity ($q = 1$) was 1.89 ± 0.38 and reciprocal Simpson’s diversity ($q = 2$) was 1.35 ± 0.21. These asymptotic values for estimated richness and diversity were reached by sampling a minimum of 11 populations (Fig. 2A). We found that total estimated richness ($q = 0$) for mycorrhizal fungi associated with A. sandvicensis by island was 3.75 ± 1.64, whereas exponential Shannon’s diversity ($q = 1$) was 3.05 ± 0.88 and reciprocal Simpson’s diversity ($q = 2$) was 2.13 ± 0.6. These asymptotic values for estimated richness and diversity were reached by sampling a minimum of two islands (Fig. 2B). Based on Mantel tests, we found no correlation between the geographic distances among populations of A. sandvicensis and the community composition of mycorrhizal fungi ($P = 0.322$), or the presence of particular Ceratobasidium OTUs ($P = 0.344$).

**DISCUSSION**

The colonization of distant oceanic islands should favour species with the ability for long-distance dispersal (Carlquist, 1974). However, for symbiotic organisms, dispersal ability alone will not determine their geographic ranges. For horizontally transmitted symbioses to establish in new habitats such as islands, either the host and symbiont(s) must arrive within similar time frames (co-dispersal) or the host must encounter compatible symbionts that are already established (co-opting). Co-opting seems the most likely strategy for plants colonizing the most remote island locations such as Hawai‘i, as it should theoretically increase chances of survival as long as a host is compatible with a wide range of local symbionts (i.e. a generalist). Conversely, if a host’s compatible symbiont(s) are few, but geographically widespread and abundant, this could also lead to successful island colonization. This latter scenario is likely to be the case for A. sandvicensis.

Contrary to our first hypothesis, we found that the obligate symbiotic orchid species A. sandvicensis is not a generalist, but that it partners with specific fungi in the genus Ceratobasidium. We circumscribed three fungal OTUs in this genus as the dominant associates of A. sandvicensis, and via phylogenetic reconstruction determined that they are closely related to taxa found throughout the globe (Table 1; Figs 3 and 4). Our most abundant and widespread Ceratobasidium taxon falls in a clade containing orchid mycorrhizal and non-mycorrhizal taxa from all the major continents except Antarctica, as well as an orchid mycorrhizal taxon from the Azores archipelago (Fig. 4). In addition to finding this taxon associating with A. sandvicensis populations across three distant and distinct islands that vary in age, size and habitats, we have also detected it in air samples from the Mauna Loa Observatory located on one of the highest peaks in the Hawaiian Islands, some 36 km from the closest A. sandvicensis (L. Tipton et al., unpubl. data). Interestingly, and counter to our second hypothesis, we found no statistically significant relationship between geographic proximity of A. sandvicensis populations and the similarity of their mycorrhizal communities or presence of particular Ceratobasidium OTUs. Combined, these results provide rare direct and indirect evidence of long-distance dispersal in fungi and insights into the ubiquity of Ceratobasidium sp. 1 across the Hawaiian Islands.

Based on our estimates of fungal diversity, we found that populations of A. sandvicensis are predicted to harbour on average between one and four fungal taxa, while individual islands are estimated to have between two and four A. sandvicensis fungal
associates present. Furthermore, our sampling efforts exceed the number of populations or islands necessary to reach community saturation for each of our estimators and have therefore probably identified the total diversity of fungi associated with *A. sandvicensis* (Fig. 2A, B). In the populations of *A. sandvicensis* where other potentially mycorrhizal lineages were detected (ex. Tulasnellaceae sp. from Oʻahu and Atheliaceae sp. from Kauaʻi, Table 1), they were in very low abundance and there is only prior evidence of Tulasnellaceae containing orchid mycorrhizal species (Dearnaley et al., 2016). While we included these lineages in our estimates of fungal diversity associating with *A. sandvicensis*, our estimates of Simpson’s diversity (approx. 1 taxon per population and approx. 2 per island) are likely to be the most realistic, as this index is weighted by the dominant taxa (Fig. 2A, B). These findings indicate that the ability of *A. sandvicensis* to establish locally is at least in part limited by its high degree of mycorrhizal specificity. In light of the high degree of mycorrhizal specificity found in
adult *A. sandvicensis* populations and the fact that many of Hawai‘i’s endemic plant species, including orchids, face extinction, below-ground symbiotic partnerships should be taken into consideration in biological conservation and restoration efforts. Additional studies that test specificity at the germination stage in *A. sandvicensis*, which may be less than adult associations (Bidartondo and Read, 2008), as well as studies of the mycorrhizal associations of the other two endemic Hawaiian orchids are needed to understand native Hawaiian orchid recruitment in the context of their mycorrhizal associations.

Mycorrhizal specificity among continental endemics and more geographically widespread orchid species may occur due to factors different from those for island endemics such as *A. sandvicensis*. Based on classic Island Biogeography Theory, the species pool of fungal symbionts should be greater on continents than on highly isolated islands such as Hawai‘i.
(MacArthur and Wilson, 2001). Therefore, the highly specific mycorrhizal associations of *A. sandvicensis* may be driven by the limited availability of compatible fungal symbionts rather than factors such as partner sanctions (Kiers et al., 2011). Both Island Biogeographic Theory and Carlquist’s ideas regarding island colonization predict that abiotic filters (e.g. distance from regional species pools) shapes island biodiversity, but Carlquist (1974) expanded upon the basic theory of MacArthur and Wilson (2001) by adding biotic filters (e.g. symbiotic interactions) into his predictions. To develop both theories further, we suggest that the spatial distribution and abundance of symbionts in addition to island age, size and isolation should also be taken into consideration for predictions of island biodiversity. Our results also offer new insights into tropical orchid mycorrhizal associations in general. Unlike other tropical orchid species, including relatives of *A. sandvicensis*, this species does not appear to be a mycorrhizal generalist-favouring Tulasnellaceae species, but forms highly specific partnerships with closely related taxa in the genus *Ceratobasidium*. Interestingly, Serendipitaceae spp. were absent from this study, providing additional support that they may be less common in the tropics (Jacqueymn et al., 2017). The degree of mycorrhizal specificity we observed in *A. sandvicensis* is similar to that of the fully mycoheterotrophic orchid species *Corallorhiza maculata* and *C. mertensiana* where populations partner with single taxa of ectomycorrhizal fungi that are closely related (Taylor and Bruns, 1999). Based on the results of prior studies, there is evidence that *A. sandvicensis* is partially mycoheterotrophic (Hynson, 2016), and that differences in the degree of partial mycoheterotrophy among *A. sandvicensis* populations may be owed to differences in fungal partnerships (Hynson et al., 2015). The current study also provides new evidence of a partially mycoheterotrophic species that forms relatively specialized mycorrhizal associations, which has previously only been found in a handful of other temperate ectomycorrhizal orchidaceous and ericaceous species (Taylor and Bruns, 1997, 1999; McCormick et al., 2004; Girlanda et al., 2005; Matsuda et al., 2012; Hynson et al., 2015).

Conclusion

In summary, by examining an endemic tropical island species that is obligately symbiotic, new patterns have emerged. First, island colonization and microbial symbiont specificity are not mutually exclusive; secondly, symbiont abundance and distribution is likely a key factor in determining the distribution of their hosts; and thirdly, in the case of *A. sandvicensis* and possibly other island-inhabiting plant species, mycorrhizal specificity may be driven by a limited available pool of compatible fungal symbionts (i.e. island host plants are occupying only a portion of their fundamental niche). While fine-level symbiont specificity, such as that observed in *A. sandvicensis*, may somewhat limit host distributions, it is not surprising that an island endemic species partners with cosmopolitan symbionts. From the host’s perspective, this strategy makes sense, as it should increase host fitness over evolutionary time by increasing the probability that it will encounter a compatible symbiont. Here we highlight how new empirical data can challenge long-standing theories of island colonization, establishment and survival, at least in Hawai‘i. In light of which, additional investigations of tropical island biogeography and biodiversity are desperately needed, especially for symbiotic organisms.

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