1	Superior dispersal ability leads to persistent ecological dominance by Candida
2	pseudoglaebosa in the Sarracenia purpurea fungal metacommunity
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12	
13	Abstract
14	Dominant taxa often emerge during microbial succession. It is not always clear what
15	makes a taxon dominant, and dispersal ability may be an overlooked mechanism
16	promoting dominance late in succession. We investigated mechanisms leading to
17	dominance of the yeast Candida pseudoglaebosa in the model carnivorous pitcher
18	plant (Sarracenia purpurea) microbial metacommunity. C. pseudoglaebosa was the
19	dominant taxon in the metacommunity, but not every individual pitcher community,
20	throughout succession. Its frequency in the metacommunity increased between early
21	and late-successional stages, and it was not replaced by other taxa. Despite its
22	dominance, C. pseudoglaebosa was not a superior competitor. Instead, it was a
23	superior disperser: it arrived in pitchers earlier, and dispersed into more pitchers, than
24	other taxa. We attribute the continuous dominance of C. pseudoglaebosa in S.
25	purpurea pitchers to its dispersal ability and the spatial structure of the

26 metacommunity. Dispersal ability influences dominance throughout microbial

27 succession.

28

# 29 Introduction

30	Primary 1	microbial	succession	occurs w	when a r	nicrobial	community c	olonizes
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31 and develops on a newly available substrate (Fierer et al. 2010). The advent of high-

32 throughput sequencing has revolutionized observational studies of microbial

33 succession, enabling researchers to describe changes in microbial communities in fine

detail (Copeland et al. 2015, Boynton & Greig 2016, Koenig et al. 2011, Cutler et al.

35 2014). These observational studies have described a variety of successional patterns.

36 For example, taxon diversity can increase, decrease, or randomly vary with

37 successional time (Cutler *et al.* 2014, Zumsteg *et al.* 2012, Redford & Fierer 2009).

38 Researchers also often observe replacement of early-successional taxa by late-

39 successional taxa with time (e.g., Boynton & Greig 2016, Wolfe et al. 2014). An

40 ongoing challenge in microbial ecology is to connect observed ecological patterns to

41 the ecological processes responsible for the patterns.

One frequently observed pattern in microbial systems is the development of
ecological dominance over successional time (Boynton & Greig, 2016, Findley *et al.*2013, Jiao *et al.* 2016). Ecological dominance is a pattern where one or a few species
comprise most of the individuals or biomass in a community (Hillebrand *et al.* 2008).
Environmental filtering, superior competitive ability of the dominant taxon, and
ecosystem engineering can all lead to dominance (Wolfe *et al.* 2014, Albergaria *et al.*2010, Goddard 2008, Nissen & Arneborg 2003, Williams *et al.* 2015). However,

49 dominance in microbial succession has been primarily studied in domesticated

50 systems, and the most frequently reported mechanisms may not be responsible for

51	dominance in all, or even most, natural microbial communities. For example,
52	environmental filtering and competition may be more important in systems where
53	human beings have designed environments to favor domesticated microbes (Wolfe &
54	Dutton 2015), and less important in natural environments with heterogeneous
55	environmental conditions.
56	Dispersal may contribute to microbial dominance over succession. When
57	primary succession happens on a sterile substrate, all members of the microbial
58	community must first disperse onto the new substrate before establishing in the
59	community. A good disperser may prevent the establishment of other community
60	members through priority effects if it arrives in a habitat first, either by pre-empting
61	or modifying available niches (Fukami 2015). Additionally, similar nearby
62	communities can impact a microbial community during succession by producing
63	propagules that then disperse into the developing community.
64	Naturally succeeding microbial communities are often physically structured in
65	groups, or metacommunities, where individual communities are spatially isolated
66	from one another but linked through dispersal (Logue et al. 2011, Holyoak et al.
67	2005). Community assembly in metacommunities is a function of ecological
68	processes occurring inside each community and dispersal among communities.
69	Ecological theory explains how dispersal can interact with intracommunity processes
70	to maintain metacommunity diversity (Leibold et al. 2004, Holyoak et al. 2005). For
71	example, populations occupying low-quality environmental patches can be
72	maintained by dispersal from high-quality patches ("mass effects"), or fitness trade-
73	offs between competitive ability and dispersal ability can maintain species diversity
74	("patch dynamics"). Theory predicts that dispersal and competition interact during
75	succession in the individual component communities of a metacommunity to result in

76 a hump-shaped relationship between successional time and species richness (Moquet 77 et al. 2003, Sferra et al. 2017). Species richness is predicted to be low early in a component community's age, to increase as more species disperse into the 78 79 community, and then to decrease as competitive interactions remove species from the 80 community. It is also possible that a particularly highly dispersing species becomes 81 dominant (and therefore decreases community diversity) in a metacommunity solely 82 as a result of its dispersal ability. Dispersal contrasts with other mechanisms leading 83 to dominance because it may allow an apparently inferior competitor to become 84 dominant through high dispersal ability. 85 We investigated the contribution of dispersal ability to ecological dominance 86 over the course of a natural microbial succession in the carnivorous pitcher plant 87 Sarracenia purpurea. S. purpurea is a perennial plant native to bogs and savannas in 88 northern and eastern North America (Buckley et al. 2003). It produces modified vase-89 shaped leaves, or pitchers (Figure 1a), annually. Developing pitchers are first entirely 90 closed, sterile chambers (Peterson et al. 2008). Once mature, the top portion of each 91 pitcher opens, and open pitchers accumulate rainwater to form small pools of water 92 (phytotelmata). Potential prey (ants and other small insects) are attracted to pitchers 93 (Bennett & Ellison 2009); some prey fall into pitchers and drown, and are then 94 shredded, decomposed, and mineralized by a food web of microorganisms and 95 invertebrates (reviewed in Ellison et al. 2003). The pitcher microbial community 96 includes bacteria, algae, and fungi, including culturable veasts (Cochran-Stafira & 97 von Ende 1998).



Figure 1: Example *S. purpurea* plant and study location. (A) One of the study pitcher plants at the edge of a *Sphagnum* island. This photograph was taken early in the growth season and both opened and unopened pitchers are visible. (B) The location of Harvard Pond in Massachusetts, USA. (C) Locations of the 43 pitchers sampled for this study. Each white circle represents one pitcher. Note that some pitchers were close enough to one another that the white circles overlap. Maps were created using ArcMap<sup>™</sup> Version 9.2 (ESRI 2006); map data are from the Office of Geographic Information, Commonwealth of Massachusetts Information Technology Division (2000) and the National Atlas of the United States (2006).

98	We followed pitcher fungal succession in a Sphagnum bog in central
99	Massachusetts where the pitcher growing season lasts for about three months
100	(although pitchers persist longer and can overwinter, Judd 1959). A single yeast
101	taxon, Candida pseudoglaebosa, was numerically dominant in the metacommunity
102	throughout pitcher growth, and its frequency increased between early and later-
103	successional stages. We used further observations and experiments to investigate the
104	ecological processes leading to C. pseudoglaebosa dominance. Unlike dominant
105	yeasts in many other systems, C. pseudoglaebosa is not an especially good competitor
106	against other tested yeasts. However, it is an especially good disperser. It is one of the
107	first fungi to arrive in pitchers and maintains high frequency in the pitcher plant
108	metacommunity over the course of the season, even as its frequency varies among
109	pitchers from completely absent to over 90% of sequences. The superior dispersal
110	ability of C. pseudoglaebosa leads it to dominate the pitcher fungal metacommunity,
111	demonstrating that dispersal ability, like competitive ability, is an important
112	contributor to ecological dominance.

#### 113

## 114 **Results**

115 Fungal communities in pitchers change over successional time

116 To understand how fungal communities develop in pitchers, we first 117 sequenced entire fungal communities in seventeen pitchers over the course of a 118 growing season. Before selecting pitchers to track for sequencing, we identified 43 119 unopened pitchers on Sphagnum islands in Harvard Pond, located in Petersham, 120 Massachusetts (Figure 1). We recorded the opening date of each pitcher and collected 121 water from each pitcher four days, seven days ("one week"), 34-42 days ("one 122 month"), and 66-74 days ("two months") after opening. At the two month time point, 123 insect herbivores, including moth larvae (Atwater et al. 2006), had destroyed ten of 124 the original 43 pitchers, and we could only sample water from 33 pitchers. In the 125 sampled pitchers, the presence of fungal DNA was assayed using the ITS1F/ITS4 126 primer pair (Gardes & Bruns 1993, White et al. 1990) Fungi were detectable starting 127 from the first measured timepoint (in 33% of sampled plants), and were widespread after one week, one month, and two months (in 91%, 95%, and 73% of sampled 128 129 plants, respectively). Seventeen of the pitchers contained detectable fungal DNA at 130 every time point from one week to two months. We sequenced fungal DNA from 131 these pitchers at all available time points, including four days if available, using 454 132 sequencing of PCR amplicons of the internal transcribed spacer (ITS) ribosomal 133 region.

Fungal succession varied among pitchers. While community composition changed significantly with time (Figure 2A), only a small amount of variation in community composition was due to variation in time (distance-based redundancy analysis adjusted  $R^2 = 0.03$ , F = 1.96, df = 1,27, p = 0.011). This small influence of

time on community composition was likely a result of high variation among pitchers. Succession followed two trajectories—five pitchers (hereafter referred to as "pitcher group 1") followed a different successional trajectory from the other twelve ("pitcher group 2")—and there was considerable variation among communities within each trajectory (Figure 2B). Distance among pitchers did not explain significant variation in community composition (partial mantel test of community composition on space controlling for time, mantel statistic r = -0.04, significance = 0.715).



Figure 2: Non-metric Multidimensional Scaling (NMDS) plots of pitcher plant community similarities. Similarities of community OTU compositions were calculated using the Jaccard metric (Jaccard, 1902). (A) NMDS plot with similarities among time points highlighted. Ellipses depict 95% confidence intervals of the centroid of each time point. No ellipse is depicted for four-day-old communities because only two were measured. (B) NMDS plot with individual pitchers highlighted. Lines connect measurements for each pitcher. Fungal communities in pitcher group 1 are represented with black lines and fungal communities pitcher group 2 are represented with gray lines. All points are located at the same coordinates in (A) and (B).

145	Despite the observed variation in fungal community composition, diversity
146	decreased, on average, in pitchers between four days and two months (Figure 3). To
147	determine a sample's diversity, we calculated Hill numbers of orders 0 to 2 ( $^{0}$ D to $^{2}$ D)
148	for each sample after rarefaction to 1040 sequences per sample. Hill numbers of
149	different orders give community diversity with an emphasis on rare species (low
150	orders) or common species (high orders) (Hill 1973, Chao et al. 2014). <sup>0</sup> D, <sup>1</sup> D, and <sup>2</sup> D



are equal to operational taxonomic unit (OTU) richness, the exponent of Shannon diversity, and the inverse of Simpson's index, respectively. Diversity as indicated by all calculated Hill numbers decreased between early and late timepoints: on average, <sup>0</sup>D declined significantly from 42.5 within the first week (including four day and one week timepoints) to 23.1 after two months (t=-3.6, df=27, p=0.001); <sup>1</sup>D declined from 14.2 to 5.2 (t=-3.8, df=27, p=0.0008); and <sup>2</sup>D declined from 8.9 to 3.2 (t=3.3, df=27, p=0.003).

159 C. pseudoglaebosa is the dominant fungal taxon in pitchers throughout succession 160 *C. pseudoglaebosa*, in the class Saccharomycetes, was the numerically 161 dominant taxon in the metacommunity, but was not dominant in every pitcher (Figure 162 4). In the metacommunity, C. pseudoglaebosa was more frequent than any other 163 taxon at every time point and its frequency increased between early- and late-164 successional timepoints. Its metacommunity frequency increased from 22% of total 165 sequences at four days to 58% at two months (Figure 4A). However, its frequency did 166 not increase over time in every pitcher. Instead, its within-pitcher frequency increased 167 over the season, decreased over the season, peaked midway through the season, or 168 dipped midway through the season, depending on the pitcher (Figure 4B). We cannot 169 say whether these increases or decreases in *C. pseudoglaebosa* sequence frequency 170 reflect changes in the total cell numbers because we did not measure cell numbers or 171 fungal biomass in pitchers. Pitcher group 1 never contained appreciable C. 172 pseudoglaebosa: each pitcher in group 1 contained less than 1.7% C. pseudoglaebosa 173 sequences regardless of the sampled time point (Figure 4B). 174 175 C. pseudoglaebosa is not a superior competitor, but has complex interactions with 176 other yeasts 177 To better understand how interactions with other yeasts might influence the 178 frequency of *C. pseudoglaebosa*, we grew *C. pseudoglaebosa* and other potentially 179 interacting pitcher yeasts in laboratory microcosms. We followed the strategy 180 advocated by Goldberg and Werner (1983), who suggested determining interacting 181 species' effects on one another by measuring organism performance as the number of 182 interacting individuals increases. We inoculated microcosms with all possible pairs of 183 three culturable pitcher yeasts (C. pseudoglaebosa, Rhodotorula babjevae, and 184



Figure 4: Taxon diversity in pitchers over time. Proportions are reported based on non-rarefied OTU assignments. (A) Taxon diversity in the entire bog metacommunity. Colored bars represent proportions of total sequences for each fungal class (or phylum for basal fungal lineages). The hatched area represents total *C. pseudoglaebosa* frequency for each time point. Note that *C. pseudoglaebosa* is in the class Saccharomycetes and represents over 99% of Saccharomycetes sequences at the one and two month time points. (B) *C. pseudoglaebosa* sequence frequency in individual pitcher communities. Data points for communities in the same pitcher are connected with lines. Black lines connect points for pitcher group 1 and gray lines connect points for pitcher group 2.

186 Moesziomyces aphidis). C. pseudoglaebosa represented 41%, R. babjevae represented 2%, and *M. aphidis* represented 0.06% of total sequences in the sequencing dataset. 187 188 Each low nutrient microcosm contained a focal yeast, which was inoculated as a fixed 189 number of cells, and an interactor, which was inoculated as a varying number of cells. 190 We then let the pairs of yeasts grow in the microcosms and investigated the effects of 191 interactors on each focal yeast using regressions. We evaluated interaction qualities 192 based on the direction (increasing or decreasing focal yeast yield with more 193 interacting cells) and shape of each regression (linear or polynomial), and we 194 evaluated differences between interactor yeasts based on whether adding interactor 195 yeast identity to each regression improved its fit. 196 Interactions between pitcher plant yeasts ranged from facilitation to 197 competition, depending on the identities of the yeasts and the number of interactor 198 cells present. Under microcosm conditions, interactions between focal yeasts and

199 interactors were polynomial when the focal yeast was *C. pseudoglaebosa* or *M.* 

200 *aphidis* (Figures 5A, B, Supplemental tables 1-4): both yeasts were facilitated by



Figure 5: Influence of interacting species on (A) *C. pseudoglaebosa*, (B) *M. aphidis*, and (C) *R. babjevae*. The plots depict the yield of each focal species as a function of the number of cells of an interacting species co-inoculated with the focal species. Interacting species are coded by color: red = C. *pseudoglaebosa*, yellow = *M. aphidis*, and blue = *R. babjevae*.

small numbers of co-inoculated cells, but their growth was impeded by larger

202 numbers of co-inoculated cells. Note that we detected facilitation of *C*.

- 203 *pseudoglaebosa* by *R. babjevae* when few *R. babjevae* cells were inoculated, but we
- 204 did not inoculate *M. aphidis* in small enough numbers to confirm *M. aphidis*
- 205 facilitation of *C. pseudoglaebosa* (Figure 5A). At high numbers of co-inoculated cells,
- 206 *M. aphidis* had a more detrimental impact on *C. pseudoglaebosa* than *R. babjevae*
- 207 (F=6.79, df=1,50, p = 0.012, Supplemental table 2). In contrast, the two interactors of
- 208 *M. aphidis* had similar effects on its yield: at low and intermediate inoculum sizes,

209 both *R. babjevae* and *C. pseudoglaebosa* promoted *M. aphidis* growth, but at high

- 210 inoculum sizes, both interactors inhibited *M. aphidis* growth (Figure 5B). Interactions
- 211 between *R. babjevae* and interactor yeasts were linear (Figure 5C, Supplemental

tables 5-6): *R. babjevae* yield was impeded by interactors regardless of the number of

- 213 interactor cells present, and *C. pseudoglaebosa* had a more detrimental impact on *R*.
- 214 *babjevae* than *M. aphidis* did (F=86.07, df=1,58, p < 0.001, Supplemental table 6).
- 215

# 216 C. pseudoglaebosa is an early disperser in pitchers

217 To investigate whether dispersal influences C. pseudoglaebosa dominance in 218 pitchers, we observed the arrival times each of the three yeasts mentioned above in 219 pitchers over the S. purpurea growth season in Harvard Pond. We surveyed the 220 presence or absence of each yeast in each of the 43 sampled pitchers using taxon-221 specific PCR primers (Table 1) to determine when each yeast arrived in a pitcher and 222 whether it persisted throughout the season. The three yeasts appeared in pitchers 223 sequentially (Figure 6): C. pseudoglaebosa first arrived in pitchers within four days 224 after the pitchers opened; R. babjevae arrived between four days and one week after 225 pitchers opened; and *M. aphidis* arrived one week to one month after pitchers opened.

- 226 Once a yeast colonized a pitcher, it either persisted in or disappeared from that pitcher
- 227 later in the season, but it did not disappear from the broader metacommunity.

228

Table 1: Taxon-specific PCR primer sequences used to detect pitcher yeasts

yeast	forward sequence	Tm	reverse sequence	Tm	product length
C. glaebosa	5'- CTGCGGAAGGA TCATTACAGT-3'	54.6	5'- TGTTCAGACAA CACTGTTCA-3'	51.8	466
R. glutinis	5'- AAGTCGTAACA AGGTTTCCG-3'	52.8	5'- CCCAACTCGGC TCTAGTAAA-3'	53.9	527
P. aphidis	5'- GGTAATGCGGT CGTCTAAAA-3'	52.6	5'- CTCTTCCAAAG AAGCGAGG-3'	53.1	467

230



Figure 6: Presences and absences of each of three yeasts in 43 pitchers over time. Each large gray shape represents the Harvard Pond at one of four time points. Circles represent pitchers: completely white circles represent pitchers in which none of the three yeasts were detected, and circles containing colored pie slices represent pitchers in which one or more of the three assayed yeasts were detected. Pie slices are colored by detected yeast: red = *C. pseudoglaebosa*, yellow = *M. aphidis*, and blue = *R. babjevae*. Circles indicate the approximate locations of pitchers, and are offset to make all data visible; see Figure 1C for accurate pitcher locations.

### 231 Discussion

#### 232 C. pseudoglaebosa is a pitcher metacommunity dominant

233 C. pseudoglaebosa was the dominant fungal taxon throughout succession in 234 the S. purpurea pitcher metacommunity, although it did not dominate every individual 235 pitcher community. It was the most frequent taxon in the metacommunity at every 236 sampled timepoint, and its frequency in the metacommunity increased after initial 237 colonization (Figure 4). Dominance in the pitcher metacommunity was in part a result 238 of the metacommunity structure itself: while C. pseudoglaebosa was not dominant in 239 every pitcher, it was dominant in enough pitchers (in some cases with a frequency 240 above 90% of all OTUs) to dominate the metacommunity as a whole (Figure 4). The 241 poor competitive performance of C. pseudoglaebosa relative to other yeasts in 242 microcosms suggests that overall competitive superiority was not the cause of its 243 dominance (Figure 5). Instead, *C. pseudoglaebosa*'s early dispersal is a more likely 244 cause (Figure 6). 245 Our best explanation for *C. pseudoglaebosa*'s dominance throughout

succession in the pitcher metacommunity is that it pre-empted other community

247 members by dispersing and establishing in pitchers before other taxa could. Early

248 dispersal likely gave C. pseudoglaebosa a numerical advantage by providing the

249 opportunity to begin exponential growth before other members of the fungal

community could arrive and begin growing (Scheffer et al. 2017). Additionally, once

251 *C. pseudoglaebosa* was established in a pitcher, facilitation by low-frequency

252 interacting taxa may have helped it maintain dominance (Figure 5A). Once

established early in the metacommunity, C. pseudoglaebosa continued to disperse into

new pitchers until between one week and one month into the approximately two

255 month long growing season (Figure 6).

256	Although C. pseudoglaebosa was the numerically dominant fungal taxon in
257	the metacommunity of pitchers, chance events, dispersal, and interactions among
258	fungi determined whether it is the dominant taxon inside any given pitcher. We expect
259	that interactions with taxa that arrive, by chance, at different times in different
260	pitchers caused the variety of C. pseudoglaebosa relative frequency changes observed
261	in pitchers ( <i>i.e.</i> , increasing, decreasing, or nonmonotonic, Figure 4B) because
262	interacting yeasts have qualitatively different impacts on C. pseudoglaebosa
263	performance depending on the number of interacting cells present (Figure 5).
264	Individual pitchers experienced priority effects because the timing of taxon arrival in
265	each pitcher (e.g., early arrival of C. pseudoglaebosa in a pitcher) determined later
266	community composition (e.g., C. pseudoglaebosa dominance in the pitcher) (Fukami
267	2015). Despite the influence of chance events on C. pseudoglaebosa dominance in
268	any given pitcher, the early and frequent dispersal of C. pseudoglaebosa compared to
269	other yeasts enabled its overall dominance in the metacommunity (Figure 4A, 6).
270	
271	Ecological patterns and processes during pitcher metacommunity succession
272	In the metacommunity, and in many individual pitchers, C. pseudoglaebosa
273	remained dominant through decreases in fungal taxon richness and diversity (Figures
274	3, 4). We did not observe a hump-shaped relationship between pitcher age and species
275	richness, as previously predicted (Figure 3A) (Moquet et al. 2003). Species richness
276	instead decreased, even as R. babjevae and M. aphidis were first dispersing into
277	pitchers late in the season (Figures 3A, 6). It is likely that C. pseudoglaebosa
278	repression of taxa through priority effects has a larger influence on species richness

than increasing diversity through new dispersal does.

280 Previous studies have also documented biotic and abiotic successional changes 281 in pitchers; while we did not measure the same parameters as these previous studies, 282 we assume that similar changes occurred in our pitcher metacommunity and that C. 283 pseudoglaebosa maintained its dominance through these changes. For example, 284 previous studies documented decreasing pH with increasing pitcher age, an early peak 285 in prey insect capture during pitchers' life spans, and a variety of changes in bacterial, 286 protist, and invertebrate community compositions over time (Fish & Hall 1978, Gray 287 et al. 2012, Miller & terHorst 2012). In bacterial, protist, and invertebrate 288 communities, the identities of dominant taxa changed as succession progressed. In 289 contrast, C. pseudoglaebosa remained the dominant fungus throughout succession. C. 290 pseudoglaebosa appears to be a classical early-successional taxon (Connell & Slatyer 291 1977, Fierer et al. 2010) because it disperses early and frequently (Figure 6), but it is 292 not replaced by late-successional taxa. 293 C. pseudoglaebosa dominance throughout succession may be enabled by the 294 short lifespans of pitchers in Harvard Pond; *i.e.*, pitchers may not live long enough to 295 enable late-succession fungal taxa to dominate the metacommunity. We sampled 296 pitchers that were up to 66-74 days old, and stopped sampling at this age because 23% 297 of pitchers had been destroyed by moths. However, pitchers in northern S. purpurea 298 populations can survive intact through winter conditions (Judd 1959), and pitchers can 299 be active for over a year in the southern United States (Miller & terHorst 2012). We 300 speculate that fungal succession more closely resembles classical successional 301 patterns and the patterns observed for other pitcher guilds (e.g., bacteria, 302 invertebrates) in longer-lived pitchers. For example, it is possible that a strong 303 competitor such as *M. aphidis* could replace *C. pseudoglaebosa* in southern *S.* 304 *purpurea* metacommunities where pitchers are active for many months. However,

305 consistent dominance of a single taxon over succession may be common in microbial
306 habitats that, like northern *S. purpurea* pitchers, have short lifespans but repeatedly
307 become available.

308

# 309 Conclusions

310	In the model pitcher plant metacommunity, taxon dispersal ability has a
311	profound influence on community structure. In particular, C. pseudoglaebosa's ability
312	to disperse into pitchers before other fungal taxa enables it to persist as the dominant
313	taxon in the pitcher metacommunity, even as intertaxon interactions and the
314	stochasticity of individual dispersal events prevent its dominance in many pitchers. It
315	is likely that dispersal ability leads to persistent dominance in a variety of other
316	natural succeeding microbial communities and metacommunities, especially when
317	early dispersal allows a taxon to prevent establishment of other taxa.
318	Future studies of microbial succession should explicitly include
319	metacommunity structure when investigating ecological processes. In the pitcher
320	metacommunity, overall taxon composition changed little over time, with C.
321	pseudoglaebosa dominant throughout succession (Figure 4A). However, individual
322	pitchers followed a variety of trajectories (Figures 2, 4B, 6). Studies of succession that
323	do not take a metacommunity's structure into account may miss community
324	heterogeneity and the diversity of ecological processes, especially dispersal ability, in
325	play among communities.
326	
327	Materials and Methods

328 Study site and field collections

329	Observations were made on Sphagnum islands in Harvard Pond, adjacent to
330	Tom Swamp, a 50 ha Sphagnum bog located in Petersham, Massachusetts at 42°30'N,
331	72°12'W (Figure 1; Swan & Gill 2007). The C. pseudoglaebosa and M. aphidis
332	isolates used in the microcosm study were collected from pitchers in Harvard Pond,
333	and the <i>R. babjevae</i> isolate was collected from a pitcher in Swift River Bog, a 2 ha
334	kettlehole bog located 75 km south of Tom Swamp in Belchertown, MA at 42°16'N,
335	72°20'W (Ellison et al. 2002). These three yeast isolates were collected in the
336	summer of 2006 and identified by comparing their ribosomal sequences, amplified
337	using the ITS1F/ITS4 and LS1/LR5 primer pairs (Gardes & Bruns 1993, White et al.
338	1990, Hausner et al. 1993 Vilgalys & Hester 1990), to sequences in the NCBI BLAST
339	database (Zhang et al. 2000). We chose C. pseudoglaebosa, R. babjevae, and M.
340	aphidis in part because they were all easily cultured from pitchers, and in part because
341	they formed colonies with different morphologies on agar plates: C. pseudoglaebosa
342	forms smooth white colonies; <i>M. aphidis</i> forms wavy white colonies; and <i>R. babjevae</i>
343	forms smooth pink colonies.
344	All S. purpurea pitcher water samples for PCR and 454 sequencing were
345	collected in the spring and summer of 2009. In May of 2009, we identified 43

346 unopened *S. purpurea* pitchers on 32 *Sphagnum* islands in Harvard Pond. Pitchers

ranged from less than 1 m to 908 m in distance to other pitchers (Figure 1C). We

348 visited each pitcher daily until it opened, and counted pitcher age from the date it

349 opened. For each pitcher water collection, the water inside a pitcher was first mixed

by pipetting up and down with a sterile plastic transfer pipette. We then removed

about 0.25 ml pitcher water and mixed it with 0.25 ml of CTAB buffer (100 mM Tris

352 pH 8.0, 1.4 M sodium chloride, 20 mM EDTA, 2% CTAB). To the best of our ability,

353 we avoided collecting insect prey or macrofauna in these samples, although any

protists and microscopic animals present in our samples were included; collected
pitcher water contained no large animal parts and appeared as a cloudy liquid. All
samples were flash-frozen in liquid nitrogen within five hours of collection and stored
at -20 or -80°C before DNA extraction.

358

359 PCR assay

We assayed each pitcher water sample for amplifiable DNA from all fungi, using the ITS1F/ITS4 primer pair, and for each of the three yeasts in the microcosm experiment, using the primers in Table 1. Primers to selectively amplify portions of each microcosm yeast's ITS sequence were designed using the NCBI BLAST primer tool (Rozen & Skaletzky 2000). We chose primer sequences to reliably amplify as much of the ITS sequence of each yeast species as possible, while not amplifying other sequences in the BLAST database.

367 To extract DNA from each pitcher water sample before the PCR assay, we 368 first thawed and centrifuged frozen samples at 16.1 g for 10 min and removed the 369 supernatant from each pellet. We then suspended each pellet in 200 µL of breaking 370 buffer (2% Triton X-100, 1% sodium dodecyl sulfate, 100 mM sodium chloride, 10 371 mM Tris, and 1 mM EDTA) (Hoffman 1997). We mixed each suspension with about 372 200 µL of 0.5 mm glass beads and 200 µL 25:24:1 chloroform:phenol:isoamyl alcohol. We vortexed each mixture for 2 min, and then centrifuged it for 5 min at 16.1 373 374 g. After centrifugation, we removed the aqueous layer and mixed it with 2.5 volumes 375 of 95% ethanol and 0.1 volume of 3M sodium acetate (Sambrook & Russell 2001); 376 we incubated each aqueous layer mixture at -20°C for at least three hours. Next, we 377 centrifuged each aqueous layer mixture for 15 min at 16.1 g, and removed the 378 supernatant. Finally, we washed each pellet with 0.5 ml 70% ethanol, centrifuged

each mixture for 10 min at 16.1 g, removed the supernatant, and resuspended each
pellet in 50 µl water.

381	We then assayed each DNA extract for the presence of each Fungal taxon, or
382	any Fungal DNA in the case of the ITS1F/ITS4 primer pair, using PCR. Each PCR
383	reaction was composed of 7.9 $\mu$ L water, 0.1 $\mu$ L GoTaq® Flexi polymerase (Promega),
384	5 $\mu L$ Flexi buffer with green dye added, 5 $\mu L$ 5x CES (combinatorial PCR enhancer
385	solution: 2.7 M betaine, 6.7 mM dithiothreitol, 6.7% dimethyl sulfoxide, 55 $\mu$ g/mL
386	bovine serum albumin) (Ralser et al. 2006), 5 µL nucleotide mix, 2 µL magnesium
387	chloride, 1 $\mu$ L of 10 $\mu$ M of each primer, and 1 $\mu$ L undiluted template DNA extract.
388	All reactions were cycled on a Biorad iCycler or myCycler using denaturing,
389	annealing, and extension temperatures of 95, 55, and 72 °C, respectively. We
390	denatured for 85 s, then ran 13 cycles of 35 s denaturing, 55 s annealing, and 45 s
391	extension, followed by 13 cycles that were identical but had a 2 min extension, and
392	finally 9 cycles with a 3 min extension. We ran a subsequent 10 min extension. Two
393	$\mu L$ of each PCR product were visualized on 1% agarose gels stained with SYBR Safe
394	dye (Invitrogen) and photographed using a U:genius gel documenting system
395	(Syngene) and a Stratagene transilluminator. Photographs of gels were scored for
396	presence or absence of a band. Bands that were too faint to reliably score were run a
397	second time with 6 $\mu L$ of PCR product per well. Presence of a band on a gel indicated
398	the presence of detectable fungal or yeast species DNA in a water sample.
399	To confirm that primers only amplified sequences from the target yeasts, we
400	randomly selected nine PCR products generated from the C. pseudoglaebosa and R.
401	babjevae primer pairs for sequencing. The primer pair that targets M. aphidis only
402	amplified DNA from six pitcher water extracts, and we sequenced all six PCR

403 products for this primer pair. Sequences were identical to or within one base of

- 404 cultured isolate sequences.
- 405

# 406 Pitcher water fungal DNA amplification and 454 sequencing

- 407 We extracted and amplified fungal DNA for fungal community amplicon
- 408 sequencing using the protocols described above, with the following changes. Gotaq®
- 409 Hotstart polymerase (Promega) was used instead of Flexi polymerase, and we used 50
- 410  $\mu$ M instead of 10  $\mu$ M of the reverse primer. The forward primer consisted of (in order

411 from 5' to 3') the 454 "A" primer (CCATCTCATCCCTGCGTGTCTCCGACTCAG)

- 412 concatenated with a 10-bp multiplex tag (454 Life Sciences Corporation 2009), and
- 413 ITS4; the reverse primer consisted of the 454 "B" primer

414 (CCTATCCCCTGTGTGCCTTGGCAGTCTCAG) concatenated with ITS1F.

- 415 Multiplex tags were unique to each sample. Reactions were cycled at 95 °C for 15
- 416 min; 30 cycles of 95 °C for 1 min, 51 °C for 1 min, 72 °C for 1 min; and a final
- 417 extension of 72 °C for 8 min.
- 418 Products were purified using Agencourt® *AMPure*® XP (Beckman Coulter)
- 419 and quantitated using a Qubit® dsDNA HS Assay (Invitrogen) according to the
- 420 manufacturers' instructions. We combined equimolar concentrations of the products
- 421 of each of three separate PCR reactions from each DNA extract. The sequencing pool
- 422 consisted of pooled equimolar concentrations of each pooled PCR product. The pool
- 423 was sequenced on one-eighth of a 454 Titanium sequencing run by the Duke Genome
- 424 Sequencing & Analysis Core Resource.

425

426 *454 sequence processing* 

427 We processed sequences using QIIME 1.3.0 (Caporaso et al. 2010). Low 428 quality sequences were removed and the remaining sequences were assigned to 429 mulitplex barcodes using the default quality filtering settings. Primers and barcodes 430 were trimmed from each sequence and sequences shorter than 200 and longer than 431 1000 bp were removed from each dataset. Sequences were denoised using the QIIME 432 denoiser. We reduced chimeric sequences by trimming the 5.8s and ITS1 portions 433 from all sequences using the Fungal ITS Extractor (Nilsson et al. 2010a), and only 434 analyzing the ITS2 portion. The 5.8s ribosomal region lies between the ITS1 and 435 ITS2 spacers, and is conserved among fungi relative to the spacers. We expected most 436 chimeric sequences to form in the 5.8s region and to be composed of ITS1 and ITS2 437 sequences from different templates (Nilsson et al. 2010b). We chose operational 438 taxonomic units (OTUs) using the uclust method in QIIME, at 97% similarity (Edgar 439 2010). We discarded all OTUs composed of a single sequence (singleton OTUs) 440 because we assumed that they resulted from sequencing errors. The longest sequence 441 in each remaining cluster was retained as a representative sequence. Of the 141,424 442 total sequences produced, 27,632 were discarded for having lengths less than 200 or 443 more than 1000 bp and 10,938 were discarded because they had low quality or did not 444 have a matching barcode. Sixty-six sequences were discarded because the ITS2 445 subunit could not be extracted and 139 sequences were discarded because they 446 represented singleton OTUs. In total, we retained 102,649 sequences for further 447 analysis. Each pitcher water sample produced between 253 and 4365 sequences. 448 449 Sequence taxonomy assignments

450 To assign genus-level taxonomy to OTUs, we performed a MEGAN analysis 451 of OTU representative sequences on the top ten hits from the NCBI BLAST

452	nucleotide database extracted using BLAST 2.2.25+ and default MEGAN settings
453	(Huson et al. 2011, Zhang et al. 2000). We discarded OTUs matching organisms not
454	in the kingdom Fungi (plant, animal, and protist sequences), but we assumed that
455	OTUs with no BLAST matches or matching unassigned fungal environmental
456	sequences were fungal sequences not yet identified in the NCBI database. We
457	retained unassigned OTUs for diversity measurements, but did not include them in
458	taxonomy summaries. We reviewed the taxon assignments output by MEGAN
459	manually, and filled in higher-level classifications (e.g., order or class) using Index
460	Fungorum (http://www.indexfungorum.org/) when an OTU was assigned to a genus
461	but not higher-level classifications. We attempted to assign all taxa to genera; if it was
462	not possible to assign a taxon to a genus, we assigned it to the most specific
463	taxonomic group possible. OTUs assigned to the genus Candida were further assigned
464	to the species C. pseudoglaebosa if they matched either C. pseudoglaebosa or its
465	close relative C. glaebosa in a BLAST search. When we compared these seven OTUs
466	against sequences representing the type strains of the two Candida species (accession
467	numbers KY102342.1 and KY102112.1, respectively), all seven OTUs aligned to C.
468	pseudoglaebosa better than they did C. glaebosa. In total, we detected 553 OTUs, of
469	which 379 were assigned to fungal taxa; fifteen were discarded because they matched
470	non-fungal sequences; and 159 were not assigned. Of the 379 fungal taxa in the
471	temporal data set, 52% were Basidiomycota, 43% Ascomycota, and 5% basal fungal
472	lineages.

473

# 474 *Microcosm interaction assays*

475 Interactions between yeasts were assayed in microcosms designed to mimic476 pitchers simultaneously colonized by different numbers of two yeast species. We

477 grew microcosms in low-nutrient media designed to mimic natural conditions in 478 pitchers. Microcosms contained sterile yeast extract media (YEM) composed of 1g/L yeast extract in local tap water (Cambridge, MA, USA). Tap water was used instead 479 480 of deionized water because we wanted the media to include micronutrients present in 481 local rainwater that may be important for pitcher plant yeast growth. The tap water 482 supply in Cambridge, MA, where this experiment was conducted, comes from three 483 Massachusetts reservoirs (Waldron & Bent 2001), and we expected it to have similar 484 inputs as rainwater in Harvard Pond pitchers. Each microcosm was inoculated with a 485 target yeast species and an interactor in 200 µL of liquid yeast media. Each target 486 yeast was inoculated with about 1000 cells per microcosm, and each interactor yeast 487 was inoculated at zero, low, medium, and high cell numbers (0 and approximately 488 100, 1000, and 10,000 cells). 489 Eighteen treatments of yeast mixtures were prepared, with ten replicates each,

490 for a total of 180 microcosms. Before inoculation, yeasts were grown in liquid YEM 491 for 48 hours. Inoculation sizes were measured after inoculation using counts of 492 colony-forming units (CFUs) on solid YEM (YEM plus 1.5% agar). Microcosms 493 were arranged in sterile 96-well polystyrene flat bottom cell culture plates and 494 incubated between 25 and 27 °C, shaking at 700 rpm for 48 hours. After incubation 495 32 microcosms were discarded because of suspected contamination. We diluted each 496 remaining microcosm, plated it to solid YEM, and counted CFUs on plates containing 497 at least 30 total CFUs. When no CFUs of an inoculated yeast were present on a plate, 498 we conservatively assumed that the yeast was present in the microcosm in numbers 499 just below our detection limit. We calculated total cell numbers assuming one instead 500 of zero CFUs for these yeasts absent from plates.

501

## 502 *Statistical analyses*

503 OTU datasets rarefied to 1040 sequences were used to produce Non-metric 504 Multidimensional Scaling (NMDS) plots and to compare community similarities, and 505 alpha diversity indices among pitchers. Eight samples contained fewer than 1040 506 sequences and were discarded. Proportions of samples assigned to taxonomic groups 507 were calculated based on the full non-rarefied dataset. Community similarities over 508 time were compared using partial distance-based redundancy analysis (db-RDA) of 509 Jaccard dissimilarity (Jaccard 1912) between each pair of samples with pitcher age as 510 the explanatory variable, conditioned on pitcher identity. A correlation between 511 geographic distance and community similarity was conducted using a partial Mantel 512 test conditioned on pitcher age. Hill numbers of order q=0 or 2 (<sup>q</sup>D) were calculated as  ${}^{q}D = \left(\sum_{i=1}^{S} p_{i}^{q}\right)^{1/(1-q)}$ , where S is the total number of OTUs and  $p_{i}$  is the relative 513 abundance of OTU *i*; <sup>1</sup>D was calculated as the exponent of Shannon diversity (Hill 514 515 1973, Chao et al. 2014). Changes in Hill numbers were modeled over time using repeated-measures linear models controlled for pitcher identity; <sup>1</sup>D and <sup>2</sup>D were log-516 transformed before analyses to homogenize variances among timepoints, and <sup>0</sup>D was 517 518 not transformed.

519 We modeled the impact of interactor yeasts on focal yeasts in microcosms 520 using multiple linear and polynomial regressions. Separate regressions were 521 conducted for each focal yeast. For each regression, focal yeast yield was the 522 dependent variable, and both the number of co-inoculated interactor yeast cells and 523 the identity of the interactor yeast were independent variables. We modeled both 524 linear and quadratic relationships between the number of co-inoculated interactor 525 yeast cells and the dependent variable because the relationship did not always appear 526 linear when plotted. Before constructing the regressions, we square-root-transformed

527 focal yeast yield to homogenize variances for the focal yeasts R. babjevae and C. 528 pseudoglaebosa, but left yield untransformed for the focal yeast M. aphidis. We also 529 transformed competitor inoculum size by  $\log_{10}(x+1)$  because interactor inoculum size 530 was varied on a log scale in the experiment. When comparing the influences of 531 competitor species, we randomly assigned treatments with no interacting yeast 532 inoculum to one of the two interacting yeasts. When selecting the best-fitting 533 regression model, we first established the best-fitting relationship (linear, quadratic, or 534 both) between log-transformed interactor inoculum size and focal yeast yield, and 535 then determined whether adding interactor identity or interactions between interacting 536 yeast identity and inoculum size to the model improved it. The best-fitting model was 537 the one with the lowest Akaike Information Criterion (AIC). All statistical analyses and index calculations were conducted using R version 538 539 3.3.1 (R Development Core Team, 2016) and the packages vegan, fields, nlme, and 540 GUniFrac (Chen 2018, Pinheiro et al. 2016, Oksanen et al. 2016, Nychka et al. 2016) 541 plots were made using ggplot2 (Wickham 2016)

542

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Supplemental Table 1: Model selection for the focal yeast *Candida glaebosa* (*C. glaebosa* yield square-root-transformed, and inoculum size  $log_{10}(x+1)$ -transformed):

explanatory variables	df	F	р	adj-R <sup>2</sup>	AIC
inoculum size	1, 51	79.37	< 0.001	0.60	405.64
(inoculum size) <sup>2</sup>	1, 51	111.1	< 0.001	0.68	394.09
inoculum size + $(inoculum size)^2$	2,50	54.65	< 0.001	0.67	395.97
(inoculum size) <sup>2</sup> + competitor identity	2,50	65.26	<0.001	0.71	389.34
$(\text{inoculum size})^2 + \text{competitor identity x} (\text{inoculum size})^2$	2,50	60.02	< 0.001	0.69	392.51
$(\text{inoculum size})^2$ + competitor identity + competitor identity x $(\text{inoculum size})^2$	3, 49	42.74	< 0.001	0.71	391.25

Supplemental Table 2: ANOVA table for the best-fitting model for *C. glaebosa*:

	df	Sum of Squares	Mean Squares	F	р
(inoculum size) <sup>2</sup>	1	10235.2	10235.2	123.73	< 0.001
competitor identity	1	561.4	561.4	6.79	0.012
residuals	50	4136.3	82.7		

Supplemental Table 3: Model selection for the focal yeast *Pseudozyma aphidis* (*P. aphidis* yield untransformed, and inoculum size  $log_{10}(x+1)$ -transformed):

explanatory variables	df	F	р	adj-R <sup>2</sup>	AIC
inoculum size	1, 58	5.37	0.02	0.07	1211.77
(inoculum size) <sup>2</sup>	1, 58	11.97	0.001	0.16	1205.83
inoculum size + (inoculum size) <sup>2</sup>	2, 57	10.86	<0.001	0.25	1199.71
inoculum size + $(inoculum size)^2$ + competitor identity	3, 56	7.64	< 0.001	0.25	1200.51
inoculum size + $(inoculum size)^2$ + competitor identity x inoculum size	3, 56	7.73	< 0.001	0.26	1200.28
inoculum size + $(\text{inoculum size})^2$ + competitor identity x $(\text{inoculum size})^2$	3, 56	7.68	< 0.001	0.25	1200.28

Supplemental Table 4: ANOVA table for the best-fitting model for *P. aphidis*:

	df	Sum of Squares	Mean Squares	F	р
inoculum size	1	173667504	173667504	6.6684	0.012
(inoculum size) <sup>2</sup>	1	391953317	391953317	15.0501	< 0.001
residuals	57	1484466512	26043272		

Supplemental Table 5: Model selection for the focal yeast Rhodotorula glutinis (R. glutinis yield square-root-transformed, and ind	oculum size
$\log_{10}(x+1)$ -transformed):	

explanatory variables	df	F	р	adj-R <sup>2</sup>	AIC
inoculum size	1, 59	92.25	< 0.001	0.60	542.62
(inoculum size) <sup>2</sup>	1, 59	67.64	< 0.001	0.53	553.45
inoculum size + $(inoculum size)^2$	2,58	45.56	< 0.001	0.60	544.44
inoculum size + competitor identity	2, 58	155.7	<0.001	0.84	489.12
inoculum size + competitor identity x inoculum size	2, 58	109.0	< 0.001	0.78	506.87
inoculum size + competitor identity + competitor identity x inoculum size	3, 57	102.1	< 0.001	0.83	491.06

Supplemental Table 6: ANOVA table for the best-fitting model for *R. glutinis*:

	df	Sum of Squares	Mean Squares	F	р	
inoculum size	1	36941	36941	225.28	< 0.001	
competitor identity	1	14114	14114	86.07	< 0.001	
residuals	58	9511	164			