- ¹ Competitive hierarchies, antibiosis, and the distribution of
- bacterial life history traits in a microbiome *

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Abstract Microbiome manipulation requires an understanding of how species interact within communities. Can outcomes of ecological interactions be predicted from microbial life history traits, the identity of the species, or both? We addressed these questions by study-13 ing the competitive interaction network in a community of 40 endophytic *Pseudomonas* spp. bacterial isolates from a native plant. Pairwise competition experiments revealed compet-15 itive dominance of P. fluorescens over P. syringae strains within this microbiome-derived community. P. syringae strains with higher growth rates won more contests, while P. fluo-17 rescens strains with shorter lag times and lower growth rates won more contests. Adding to their competitive dominance, P. fluorescens strains often produced antibiotics to which few 19 P. syringae strains were resistant. Many competitive outcomes among P. syringae strains 20 were predicted to be reversed by P. fluorescens inhibitors because indirect benefits accrued 21 to less competitive strains. P. fluorescens strains frequently changed competitive outcomes, 22 suggesting a critical role of strains within this bacterial clade in structuring plant microbiome communities. Microbial traits also may provide a handle for directing the outcome of colonization processes within microbiomes.

Keywords: Pseudomonas, indirect interactions, phyllosphere, microbiome, phytopathogen

Code and data available at https://github.com/phumph/competitive_hierarchies.

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27 Introduction

The ecological forces shaping bacterial microbiome community structure are difficult to characterize, given the diversity and relatively uncultivable nature of these taxa, particularly in animals. Plants, in contrast, possess a highly cultivable microbiome and have potential to serve as models for understanding microbiome ecology and evolution generally. Moreover, plant growth-promoting bacterial (PGPB) formulations are being deployed in agriculture. Quantifying and predicting ecological outcomes among common species in these artificial communities is therefore also of practical value. Competition may be the principle ecological force shaping microbial community structure 35 (Foster & Bell 2012; Coyte & Rakoff-Nahoum 2019), yet distinct forms of competition can operate within communities: competition for shared resources and interference with another 37 species' ability to do so (Case & Gilpin 1974). In addition to structuring microbiome communities, competition of both types is a potent source of natural selection (Hibbing et al. 2010; Cornforth & Foster 2013; Mitri & Foster 2013). Teasing apart how exploitative and interference competition interact in a community context remains a challenge more generally (Amarasekare 2003; Delong & Vasseur 2013; Coyte et al. 2015). Furthermore, as diversity increases, the number of possible indirect interactions in the community scales faster than the number of direct interactions. Accordingly, a species may benefit from additional competitors if the net indirect effects dampen direct competition faced by other species (Levine 1976; Lawlor 1979; Stone & Roberts 1991; Wootton 1994; Miller & Travis 1996). Such indirect facilitation has not been well explored in microbiomes. Species-rich communities are also more likely to harbor members with traits that have a large ecological impact (Banerjee et al. 2018). In microbial communities, strains that secrete diffusible antibiotics, resource substrates, or signaling molecules can alter the fitness of nonproducers (Lee et al. 2010; Gutiérrez & Garrido 2019). By selecting for more specialized traits involved in resistance or metabolite uptake, these secretions can upend competitive

hierarchies that would otherwise be mediated by canonical competitive fitness traits. It is unclear if microbial taxa with large indirect impacts are common in natural microbiomes (Banerjee et al. 2018). Leaf-dwelling (phyllosphere) bacteria secrete compounds altering growth and survival of nearby bacteria (Lindow & Brandl 2003; Quiñones et al. 2005; Dulla & Lindow 2009; Dulla et al. 2010) and can co-localize on the leaf surface and interior (Monier & Lindow 2005). Thus, there is potential for direct and indirect interactions between competing bacteria to affect community assembly and steady-state patterns of diversity in plant microbiomes.

Finally, competition need not be purely hierarchical: intransitive loops may arise in species-

Finally, competition need not be purely hierarchical: intransitive loops may arise in speciesrich communities whereby numerical dominance cycles at local spatial scales, resulting in
community stability (Kerr et al. 2002; Rojas-Echenique & Allesina 2011). Even modest intransitivity can buffer against extinction (Laird & Schamp 2006; Rojas-Echenique & Allesina
2011; Laird 2014) and the degree of intransitivity can shape species diversity (Reichenbach
et al. 2007). Although intransitivity occurs in microbial systems in the laboratory (Kerr et
al. 2002; Kelsic et al. 2015), its occurrence in natural microbiome communities is not well
understood (Lankau et al. 2011; Godoy et al. 2017).

To address the various gaps highlighted above, we studied how microbial traits mediate direct and indirect competitive outcomes in an assemblage of co-occurring bacterial species from a wild, endophytic microbiome meta-community. Specifically, we (1) characterized life history trait variances and co-variances of diverse isolates in the laboratory, (2) examined how such traits related to competitive interaction networks manifest in spatial microcosms, and (3) analyzed whether indirect interactions among strains might be expected to strengthen or weaken competitive hierarchies among strains, with the latter expected to promote coexistence under natural conditions. We used a diverse set of endophytic *Pseudomonas* spp. bacteria derived from native bittercress (Brassicaceae: *Cardamine cordifolia* A. Gray), encompassing an extensive sample of the diversity found in both the putatively phytopathogenic *P. syringae* clade and the presumed saprophyte *P. fluorescens* clade (Humphrey *et al.* 2014;

80 Humphrey & Whiteman 2020).

81 Methods

Overview

We measured a network of pairwise competitive interactions among 40 *Pseudomonas* spp.

strains, wherein strains competed for shared resources in spatial microcosms. We quantified

each strain's ability to invade and defend against invasion and derived a composite measure

of competitiveness that incorporated both invasive and defensive ability. We simultaneously

measured each strain's capacity to interfere with growth of surrounding competitors through

inhibitory secretions, as well as each strain's apparent ability to resist such inhibitors. Using

independent measurements of maximum rate of increase, lag phase, and maximum yield *in*vitro, we then determined the underlying correlates of both exploitative and interference

competitive abilities, as well as effect of phylogenetic distance on these correlations. Fi
nally, using the distribution of pairwise outcomes measured in our competition assays, we

inferred the number and direction of indirect interactions that would result in facilitation

via inhibition of a superior competitor by a nearby producer strain.

95 Bacterial strains

Of the 51 Pseudomonas spp. strains isolated from bittercress and previously described (Humphrey et al. 2014), we selected a set of 40 (26 P. syringae, 14 P. fluorescens) that represented the phylogenetic diversity present in this community (Humphrey & Whiteman 2020). We included the laboratory strain P. syringae pv. maculicola str. ES4326 in our strain set owing to its phylogenetic similarity to strains isolated from bittercress and its extensive characterization in the laboratory as a pathogen of Arabidopsis thaliana (Cui et

al. 2002, 2005; Groen et al. 2013). All bacterial strains used had undergone only one prior growth cycle after freezing following initial isolation on King's B plates from surfacesterilized homogenates of bittercress leaf samples (Humphrey et al. 2014). For each strain,
we estimated resource usage (i.e., growth) parameters (maximum growth rate r, lag phase L, maximum yield K) from in vitro growth cycles conducted in 96-well plates (see Online
Supplemental Materials [OSM]: Supplemental Methods for details).

Pairwise competition assays

We conducted pairwise high-density competition assays in spatial microcosms in which a 109 "resident" strain inoculated onto the surface of each plate competed with each "invader" 110 strain spotted on top (see **OSM**: Supplemental Methods for details). We visually scored 111 growth of each invader as 0 for no visible growth of the invader above a negative control 112 spot containing sterile growth media alone, 0.5 for a largely translucent 'megacolony', which 113 reflected a definite presence of growth but which was relatively suppressed and confined 114 to the megacolony margin, and 1 for obvious and robust megacolony growth. We scored 115 inhibition interactions as a binary outcome indicating the presence of a zone of clearance 116 (halo) ≥ 1 mm surrounding the extent of the invader megacolony. 117

118 Calculating indexes of competitiveness

Each strain was assayed under 40 different conditions both as resident strain and invader, comprising an interaction network with 1,600 entries (including self vs. self). One version of the interaction network represents the outcome of resource competition and details the extent of growth of each invader, while the other captures the presence or absence of inhibitory interactions indicated by zones of clearance in the resident population. For resource competitions, we calculate the invasive capacity (C_o) and defense capacity (i.e. territoriality; C_d) of each strain. C_o for each strain i was calculated as

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$$C_{o,i} = \frac{1}{n_{ij}} \sum_{i \neq j}^{n} x_{ij}$$

where $x_{ij} \in \{0, 0.5, 1\}$ and n_{ij} is the total number of scored interactions for each strain as the invader with all non-self resident strains. C_o is thus the expected value of growth attained by each strain as the invader across the population of residents. Similarly, C_d quantifies the ability of each strain to resist invasion by other strains and is calculated as

$$C_{d,j} = \frac{1}{n_{ji}} \sum_{j \neq i}^{n} (1 - x_{ji})$$

Here, strain j is in the resident state, and $x_{ji} \in \{0, 0.5, 1\}$ as before but with a subscript reversal, indicating the degree to which the resident prevented the growth of each invader i. As above, n_{ji} is the number of interactions occurring between each focal resident and its non-self invaders. C_d can thus be interpreted as the expected amount of growth each resident strain can prevent among the population of invaders assayed.

We then calculated an overall exploitative competition index, C_w , for each strain as

$$C_w = C_o - (1 - C_d)$$

where $-1 \le C_w \le 1$. These extremes represent absolute competitive inferiority (-1), where

a strain failed to prevent all growth of any invader and similarly failed to invade any other strain, to absolute competitive dominance (1), where a strain fully invaded all residents and fully prevented growth of all invaders. We also calculated C_t and C_r based on the interaction matrix for interference competition. Here, C_t is the proportion of successful invasions (i.e., given growth of 0.5 or above) that also resulted in halo formation produced by the invading strain, indicating inhibition of the resident. C_r for a strain is the proportion of contests with all invading inhibitor strains

(i.e., all strains with $C_t > 0$) that failed to result in halo formation, which we interpreted as resistance. Analogous to C_w above, we calculated an overall interference competition index, I_w , as

$$I_w = C_t - (1 - C_r)$$

where $-1 \le I_w \le 1$, which is analogous to the aggressiveness index (AI) of (Vetsigian *et al.* 2011).

Analyzing the distribution of competitive outcomes

We determined when outcomes of all pairwise interactions between strains i and j $(i \neq j)$ took the following forms: reciprocal invasibility (RI), where strains i and j each invade one 151 another; reciprocal non-invasibility (RNI), where strains i and j cannot invade each other; 152 and asymmetric (Asym), where strain i invades strain j but j cannot invade i. To compare 153 outcome distributions, we constructed binomial linear models to estimate the probability of 154 RI, RNI, and Asym as a function of bacterial clade (P. syringae versus P. fluorescens). 155 In addition, we compared trait co-variances and overall levels of trait dispersion between 156 P. syringae and P. fluorescens, correcting for phylogenetic distance between strains in each 157 clade. To do so, we first we conducted principal components analysis (PCA) using the matrix 158 of mean-centered and scaled competitive indexes and growth parameters for all strains (40 x 159 9 matrix) as input. We then calculated Euclidean distance between vectors of [PC1, PC2, 160 PC3] for all pairs of strains within each *Pseudomonas* clade. Using these calculated pairwise multivariate trait distances as a response variable, we computed linear regression models with bacterial clade as well as phylogenetic distance (D_g) as predictors. We calculated D_g as the pairwise uncorrected nucleotide distance between 2,690 bp of sequence comprised of four 164 partial housekeeping gene sequences previously generated for each strain from Humphrey 165

of overall competitiveness (C_w) .

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et al. (2014). Orthologous sequences from the genome of Psm4326 were derived from its published genome sequence (Baltrus *et al.* 2011); RefSeq ID NZ_AEAK00000000.1).

Inferring indirect interactions from the pairwise network

We next examined the structure of the pairwise competitive interaction network for signa-169 tures of intransitivity (i.e., non-hierarchical or context-dependent interactions). Using data 170 from pairwise interaction outcomes, we assessed (1) whether three-strain competitions would 171 result in intransitive loops (e.g., rock-paper-scissors outcomes) such that no species would be 172 globally dominant; and (2) whether the presence of secretions from a nearby P. fluorescens 173 strain would reverse the outcome of a pairwise interaction that would typically result in 174 competitive dominance of a single strain (indirect facilitation). Facilitation can occur by 175 strain A releasing strain C from inhibition from strain B (where A also has to be resistant to 176 B's inhibitors), or from resource competition from superior competitor strain B. This analysis 177 is agnostic to mechanism but calculates the proportion of conditions under which facilitation 178 of an otherwise weaker competitor is expected to arise. A total of 8,203 trios were analyzed 179 for potential facilitation based on the pairwise interaction data from 641 pairs of strains that 180 met the competitive asymmetry criteria. 181 For each strain, we calculated the net effect of antagonistic vs. facilitative indirect interactions 182 across all possible trios and compared this to underlying fitness metrics derived from the 183 pair-wise interaction network. We then compared how strongly the net effects from indirect 184 facilitation are expected to change fitness ranks of strains in relation to their baseline values 185

187 Results

188 Competitive outcomes

Pairwise soft-agar invasion assays revealed that the competitive ability of P. fluorescens 189 strains was consistently superior to P. syringae strains (Fig. 1): ~99\% of strain pairings 190 between the two clades resulted in asymmetric dominance of P. fluorescens over P. syringae 191 (99% Asym; Fig. S2; Tables S1, S2). Within P. fluorescens, the proportion of reciprocally 192 non-invasible (RNI) pairings was significantly higher compared to within P. syringae pairings 193 (Fig. S2; Tables S2, S3). The competitive dominance of P. fluorescens over P. syringae was 194 evident across both exploitative and interference-based measures of competitiveness (Figs 1, 195 2; Table S1). 196

197 Interference competition

Of the 40 strains assayed, 13 (all P. fluorescens) produced halos surrounding some subset 198 of the resident strains they invaded (antibiosis), indicating the production of antibiotics 199 (diffusible inhibitors/toxins) (Fig. 1). Mean inhibition index (I_w) among P. fluorescens 200 strains was 0.15, although two strains inhibited only one other, and P. fluorescens strain O3A failed to inhibit any strain (Fig. 1). Four P. fluorescens strains were susceptible to inhibition by two of the toxic strains (43A, 34A; Fig. 1). Resistance to toxin producers in P. syringae was variable, although the mean value was high at 0.72 (Fig. 2b; Table S1). In at least one case, resistance among P. syringae strains showed a strong correlation with 205 phylogenetic position: invading strain P. fluorescens str. 43A adopted distinctly different 206 phenotypes in pairings with P. syringae strains from different sub-clades (perMANOVA F =207 7.04, 1000 permutations, p = 0.002; Fig. S3). Nine of the 25 43A megacolonies had a smooth 208 morphology, 13 adopted a highly motile morphology we call the "smooth spreader", and 200 the three remaining adopted a wrinkly spreader-like morphology (Fig. S3a-c). Inhibitor 210

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production by 43A was strongly associated with the smooth morph ($\chi^2 = 19.2$, p < 0.001; Fig. S3e); 43A only inhibited one strain as the smooth spreader morph, and then only after it had stopped expanding across the plate (personal observation). None of the three wrinkly spreader-like morphs produced toxins that inhibited a resident strain.

The correlations between competition and growth traits showed opposite patterns for strains

Life history correlates of competitiveness

within P. syringae versus P. fluorescens: overall exploitative competitiveness (C_w) was neg-217 atively correlated with both r and L for P. fluorescens (Pearson's $\rho = -0.78, -0.75, \text{ re-}$ 218 spectively; Fig. 2c). That is, P. fluorescens strains with shorter lag (smaller L), and thus 219 smaller r, were more competitive in our assay. This apparent trade-off between maximum in220 vitro growth rate r and growth initiation (1/L) was not observed across P. syrinage strains. 221 Instead, C_w in P. syringae was positively correlated with only r ($\rho=0.78;$ Fig. 2c). Strains 222 from neither clade showed a canonical trade-off between r and in vitro saturation density 223 (K). On the contrary, P. syringae strains showed a positive correlation between K and 224 growth rate as well as defensive capacity C_d , while for P. fluorescens K was positively cor-225 related with levels of resistance (C_r) . Overall, offense (C_o) and defense (C_d) were strongly 226 positively correlated with linear slopes near 1 for both clades (Fig. 2c; Fig. S5). All three 227 measures of exploitative competition were positively related to interference measures for P. 228 fluorescens (Fig. 2c). 229 Principal component analysis (PCA) of all nine traits revealed largely non-overlapping 95% 230 confidence ellipses for the two clades (Fig. 2d). The first two PCs together explained 72.5% 231 of the variation in the data. The loading vectors of C_w and lag duration were in opposing 232 directions, indicating a negative correlation, while those for competitiveness and inhibitory 233 capacity are largely co-linear, indicating a positive correlation (Fig. 2d). The loading for 234 resistance, C_r , was nearly co-linear with lag duration, a relationship not apparent in the 235

pair-wise correlation analysis in Fig. 2c.

Overall, strains within the P. syringae clade showed greater intra-clade pairwise trait differences across PCs 1-3 than strains within P. fluorescens (Welch's unequal variants t test, t = 8.7, $p < 10^{-6}$; Fig. S7). While multivariate trait distance increased on average with phylogenetic distance (D_g term $\beta = 0.1$, $p < 10^{-10}$; Table S4), P. syringae strains showed a higher average trait distance even after accounting for D_g in a multiple regression model (Psyr term $\beta = 0.9$, $p < 10^{-8}$; Table S4).

²⁴³ Competitive interaction network and intransitivity

Five trios met the criteria for a rock-paper-scissor game out of the 9,604 possible trios of 244 interactions evaluated (Fig. 3a). Nine unique strains were implicated in these trios. Each 245 trio was comprised of distantly related P. syringae strains (mean D_G between strains in 246 R-P-S trios = 0.118 [0.115 - 0.122 95% CI]. A further 632 (7.7%) met the criteria whereby the inferior competitor was facilitated by the inhibition of the superior competitor by a third party to which the facilitated strain was resistant (Fig. 3a). Despite the overall tendency to reinforce pairwise interactions, indirect facilitation from 250 inhibitor-producing strains implicated nearly all (39) of the 40 studied strains in one or more 251 of the three possible trio roles: the facilitator, the knocked-out competitor, or the facilitated 252 strain (A, B, and C, respectively; Fig. 3a). Overall, 26 strains were facilitated (C), and 253 21 of these also served as the knocked-out competitor (B) in a subset of the trios (Fig 3b, 254 inset). Twelve of the 13 inhibitor-producing strains (all P. fluorescens) were implicated as 255 facilitators (A strains) (Fig 3b, inset). 256 Intuitively, the propensity towards B vs. C roles was correlated by underlying differences in competitive fitness: the most facilitated strains (high C fraction) were among the least competitive (low C_w) in the population, indicated by a negative correlation (r = -0.76[0.86 - 0.58] 95% CI], $p < 10^{-5}$; Fig. 2c). B strains were intermediate relative to the entire 260

range of C_w values. Facilitator A strains had consistently higher C_w , owing to the generally 261 higher competitiveness of P. fluorescens strains: in all but 6 of the 632 facilitation trios, the A strain out-competed the C strain in the pairwise network, even though such strains 263 were resistant to their inhibitors (Fig 3b). This finding suggests that facilitation in this 264 network depends on it occurring at a distance whereby the facilitator does not immediately 265 out-compete the resistant strain which it facilitated. Also intuitively, resistance (C_r) was 266 strongly positively correlated with the probability of being facilitated (Pearson's $\rho = 0.57$ 267 $[0.32 - 0.75 95\% \text{ CI}], p < 10^{-4}.$ 268 Only rarely were P. fluorescens strains anything other than the facilitator strain: only three 269 were ever knocked out by an A strain to which they lacked resistance (36A, 46A, 06B). 270 This finding reveals that P. fluorescens strains very rarely benefit from indirect facilitation, 271 in contrast to their frequent role as facilitator (Fig. S8). One strain (P. fluorescens str. 272 43A) played the role of facilitator (A) in >25% of all facilitation trios, over 2.5-fold more often than the next most frequent facilitator (Fig. 3b). This indicates that the presence 274 of individual inhibitor-producing community member can substantially shift the outcome 275 distribution among non-producers. 276 Averaged across all inhibitor-producing strains, the net effects of indirect interactions reshuf-277 fled the fitness ranks of P. syringae strains to a degree that weakens the original pairwise 278 competitive hierarchy (rank correlation ρ between C_w and C_w' = 0.50; Fig. 3e; Fig. S8), 279 such that the former advantage of several top P. syringae competitors gets redistributed 280 across a larger number of relatively weaker competitors. In contrast, the hierarchy among 281 P. fluorescens strains was generally recapitulated, or exacerbated, by indirect interactions 282 arising from antibiosis in this network. 283

Discussion

Overview Overview

We discovered clear clade- and trait-level associations with the outcomes of competitive 286 interactions among naturally occurring bacterial strains. Using a subset of endophytic bac-287 teria isolated from a native sub-alpine plant (C. cordifolia), we found major differences in 288 both exploitative and interference competitiveness between the two principle Pseudomonas 280 spp. clades in this endophytic community. Trait co-variance structure revealed the biologi-290 cal differences between these two major clades of native plant-associated *Pseudomonas* spp. 291 bacteria. Such patterns suggest that the evolution of competitiveness may involve distinct 292 components of life history in these bacterial lineages. When placed into an ecological context, 293 the trait distributions we revealed across this bacterial assemblage are predicted to generate 294 context dependence in competitive outcomes in the form of facilitation, whereby a inhibitor 295 strain displaces a strong competitor and thereby facilitates a resistant but weaker recipient. 296 Thus, the community context of interference competition is important for predicting the outcome of competitive pairings which typically depend primarily on exploitative capacity. Such a dataset allows dissection of several dimensions of in vitro fitness exhibited by a 299 natural community of phyllosphere *Pseudomonas* spp. and provides a platform for testing 300 hypotheses about the mechanistic bases of competitive traits (e.g., toxin production and 301 resistance) and their potential effects on ecological diversity and microbiome community 302 structure. We also showed that P. fluorescens, presumed to be soil dweller, can be both 303 common and important in structuring the outcome of ecological interactions within the 304 context of the leaf microbiome. Together, this work helps build an understanding of how 305 competitive traits might evolve in tandem with other life history traits in representatives 306 from real communities that interact in nature.

Correlations between growth traits and competitiveness

Neither P. syringae nor P. fluorescens strains exhibited canonical growth rate trade-offs 309 with maximum yield, K, which can result in a tragedy of the commons whereby rapid but 310 wasteful use of resources yields higher competitive ability (Pfeiffer et al. 2001; MacLean 311 2008). Rather, a more pronounced signal was that maximum growth rate was correlated 312 with a longer lag phase in P. fluorescens. This pattern contradicts the traditional dichotomy 313 between generally "fast" vs. "slow" life histories and contrasts with patterns observed in 314 microbial evolution experiments. For example, Escherichia coli lines adapting to a glucose-315 limited environment exhibited coordinated increases in growth rate and shorter lag time 316 after 10,000 generations (Vasi et al. 1994; Lenski et al. 1998). Additionally, E. coli selected 317 to persist in lag phase during periods of antibiotic stress exhibited no pleiotropic changes in 318 maximum growth rate despite up to a 10-fold increase in lag time (Fridman et al. 2014). Our 319 study adds support for the idea that lag phase deserves attention as an important feature of 320 microbial life cycles, and characterizing the physiology of cells during this phase may reveal 321 the nature of its correlations with maximum growth rate and competitive fitness in this and other systems.

The negative correlation between lag phase and growth rate in P. fluorescens resembles a 324 colonization—competition trade-off. Spatial priority effects arising from territoriality can pro-325 vide a mechanism for maintenance of colonization—competition trade-offs that would other-326 wise lead to competitive exclusion (Edwards & Schreiber 2010). A colonization—competition 327 trade-off underlies territoriality in Vibrio spp. based on the differential ability of clones to 328 contest territory vs. disperse to new ephemeral habitats (Yawata et al. 2014). One hy-329 pothesis arising from our work is that P. fluorescens strains that preempt as much space 330 as possible within patchy and ephemeral leaf environments may reap the rewards of their 331 territorial monopoly even at the expense of a decreased maximal growth rate. 332

The production of exudate (C_t) or exudate resistance (C_r) did not trade-off with any of the

life history traits we measured (Fig. 2a). This is consistent with findings that exudate production did not affect in vitro growth rates measures in P. fluorescens (Garbeya et al. 2011). Instead, we found a positive correlation between inhibitory ability (C_t) and overall exploita-336 tive competitiveness for P. fluorescens. Although perhaps unexpected from a theoretical 337 perspective (Neumann & Jetschke 2010), such a positive correlation is nevertheless intuitive: 338 megacolonies invading a resident strain presumably must reach a critical size in order for 339 any toxicity to be detectable if induction is either density dependent or if the toxic effects 340 are concentration dependent. Cells may only reach such a critical density if their relative 341 exploitative competitiveness enables them to do so, without which interference competitive 342 ability is irrelevant. Further empirical work, scaling from individual cells to populations, 343 will be required to properly ground co-existence theory for microbes in mechanistic models 344 of trait-trait interactions. 345

Our study is limited in that we relied on visible manifestation of growth inhibition. Interference mechanisms range from direct injection of bacterial effectors via Type VI Secretion Systems (Decoin et al. 2014, 2015), the production of subversive growth-regulating secreted N-acylhomoserine lactones (AHLs) or enzymes that quench these signals typically involved 340 in quorum sensing (Dulla & Lindow 2009; Dulla et al. 2010), or the production of secreted 350 toxins (e.g. bacteriocins or phage-derived proteins). Further work is needed to describe 351 the range of interference mechanisms that may operate within plant microbiomes and to 352 characterize the ecological effects of newly described modes of interference capable of being 353 deployed by P. syringae (Hockett et al. 2015; Kandel et al. 2020) that this study was not 354 capable of detecting. 355

356 Ecological implications

If strains from *P. syringae* and *P. fluorescens* were to compete in an unstructured environment, where preemption of space was irrelevant, *P. syringae* strains with high growth rates

might be expected to out compete a variety of P. fluorescens strains with relatively lower growth rates (Fig. 2). But within the structured and ephemeral context of the leaf environment, P. fluorescens may act as a territorial species whose potential effect in the phyllosphere 361 may be to exclude colonization by other strains including P. syringae. This is consistent 362 with the identity of P. fluorescens as a plant mutualist, although the evidence of this comes 363 exclusively, to our knowledge, from studies of its indirect effects via plant defensive signaling 364 or direct toxicity to pathogenic fungi following its colonizing of plant roots (Mendes et al. 365 2011: Hol et al. 2013). In addition to such indirect effects, the superior competitiveness of P. 366 fluorescens over P. syringae suggests that direct interactions may affect phyllosphere bacte-367 rial community assembly and plant disease risk from phytopathogenic isolates of P. syringae. 368 Irrespective of the underlying mechanisms of interference and resistance, the frequency of 360 these traits in a community may have large indirect effects that generate context-dependent 370 competitive asymmetries among diverse genotypes. 371

The ecological context in which traits are expressed impacts functional diversity (both genetic and phenotypic) found within natural communities (Ohgushi et al. 2012), despite 373 strong pairwise competitive asymmetries, as seen here between *Pseudomonas* spp. clades. 374 In our interaction network, indirect effects of interference competition may equalize fitness 375 differences between P. syringae competitors that otherwise have asymmetric exploitative 376 abilities (Fig. 3b; Fig. S8). Facilitation of the sort explored here is only possible with an 377 intermediate frequency of toxin resistance expressed by P. syringae (Fig. 3d). The fact that 378 resistance is not more common among P. syringae suggests a cost of resistance that did not 379 manifest itself in the assays conducted in our study. Further study into the mechanisms of 380 production of, and resistance to, interference traits in this community would help explain 381 the distribution of these traits in the community as well as their costs and correlations with 382 other traits. 383

We show that the gains from facilitation are predominantly accrued by weaker resource competitors (Fig. 3c-f; Fig. S8). Only in a small subset of the facilitation trios could the

facilitated strain invade the producer. When the facilitated strain does not pose a competitive threat to the facilitator—as is the case most of the time here—the gains from facilitation may be short-lived. However, the overall effect of this degree of facilitation may be to prolong 388 periods between exclusion/extinction events, elevating the diversity that is observable at any 389 given point within the system (Laird & Schamp 2006). The additional form of intransitivity 390 found in our study is a pair of extended trios that have R-P-S invasion asymmetries, which 391 are predicted to lead to frequency-dependent or cyclical invasion dynamics (Laird 2014). This 392 prediction is awaiting an empirical test, and this system presents an excellent opportunity 393 for doing so. 394

395 Conclusions

We found that competitive abilities of strains within a natural assemblage of plant-derived Pseudomonas spp. varied between the two major clades present, P. fluorescens and P. sy-397 ringae. Competitive fitness in our assays hinged on different traits in these two clades, and 398 the higher degree of inter-strain trait dispersion in P. syringae may indicate that the focal 399 traits measured here undergo more rapid evolution given the same degree of phylogenetic 400 divergence (Fig. 2d; Fig. S7). We found no apparent life history trade-offs between growth 401 rate and yield. Although speculative, the P. fluorescens clade may contain early colonizing 402 strains that contest territory to a greater extent, which may serve to directly buffer against leaf colonization from potentially phytopathogenic P. syringae. In contrast, a high degree of inhibitor resistance among P. syringae may prevent local exclusion when spatial structure releases them from direct exploitative competition with P. fluorescens. Finally, the combination of exploitative and interference competition due to inhibitor-mediated facilitation 407 may stabilize co-existence of strains that otherwise competitively exclude one another. Our 408 study sheds light on the types of ecological interactions between bacterial lineages within mi-409 crobiomes that should be quantified during development of microbial formations for clinical 410

and crop enhancing purposes.

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417 Competing interests

The authors declare no competing interests.

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Figure captions

Fig. 1. Pairwise competitive interactions in a phyllosphere *Pseudomonas* spp. community. Rows reflect strains in the resident state, while columns reflect strains in the invader state. Dashed red lines through interaction matrix denote within–between clade divisions for ease of visualization. Phylogeny modified from Humphrey et al. (2014).

Fig. 2. Phylogenetic distribution of life history trait variation within a Pseudomonas spp. community. a. Life history components are maximum growth rate (r_m) , lag phase (l), maximum yield (K), derived from individual microcosm growth experiments; and components of offensive (C_o) , defensive (C_d) , overall (C_w) competitiveness, resistance to toxicity (C_r) , toxicity (C_t) , and overall interference capacity (I_w) derived from a pairwise competitive interaction network (see Methods). Column z-score of each trait value indicated by color. b. Smoothed frequency distributions of trait values for each measured trait by clade (P. fluorescens) and P. syringae). c. Pairwise correlations and principle component analysis (PCA) (d) of nine traits revealed clear dissimilarities in trait distributions and patterns of co-variance between clades. Correlations with text values reflect magnitude of each Pearson's ρ where the FDR corrected p < 0.05; comparisons with FDR-corrected p < 0.10 are italicized. d. PCA 95% envelopes per clade depicted as solid or dashed ellipses. Dots are labeled with strain IDs. Individual trait vector loadings are in blue for resource use traits and orange for competitive indexes).

Fig. 3. Prevalence of intransitive interactions in a Pseudomonas spp. interaction network. a. Types of interaction trios resulting in facilitation or rock-paper-scissors (R-P-S) competitive asymmetries. N = number of trios meeting the given criteria out of the total trios analyzed (see Methods). b. Frequency distributions of how often each strain played the facilitator (A), the knocked-out intermediate (B), or the facilitated (C). Several strains

played multiple roles; strains in facilitative trios with as well as without toxic intermediates are indicated with black triangles to the left of the strain IDs. Panel (**b**) inset displays the distribution of the number of unique strains that played each combination of roles. Only 06B played all three roles. The probability of playing A, B, or C roles in facilitative trios varied with (**c**) overall competitiveness, C_w , as well as (**d**) resistance. **e.** ΔC_w plotted against baseline C_w shows initially weaker P. syringae strains benefit the most from indirect interactions (Pearson's $\rho = -0.67$), while P. fluorescens fitness remains relatively unaffected by indirect interactions ($\rho = 0.74$). **f.** Net competitive fitness ($C'_w = C_w + \Delta C_w$) after considering indirect effects weakens competitive hierarchies among P. syringae ($\rho = 0.50$) but has little effect on P. fluorescens competitive fitness ($\rho = 0.98$).

Figures

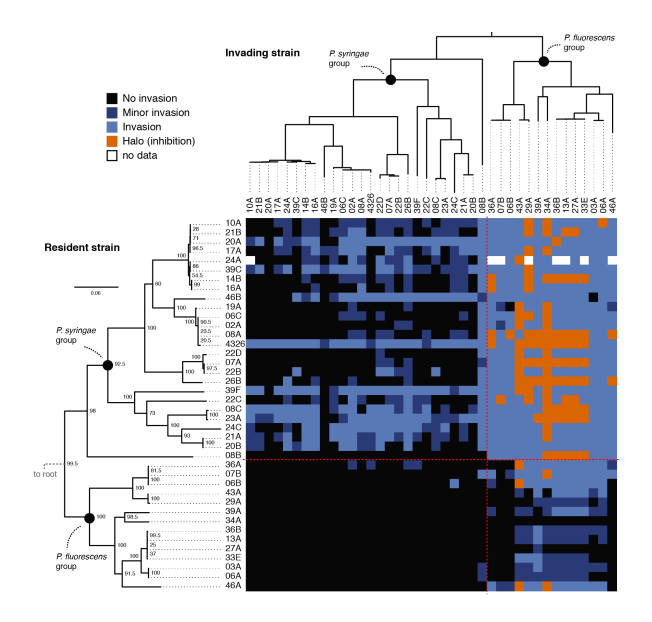


Figure 1:

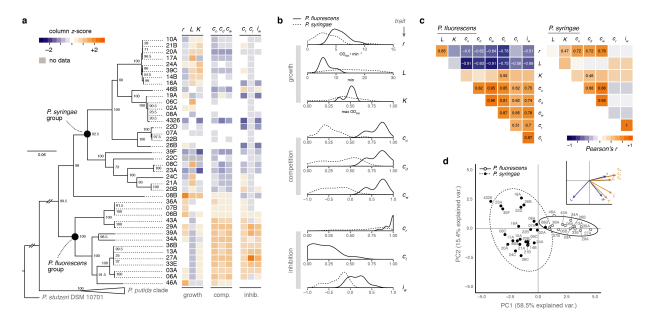


Figure 2:

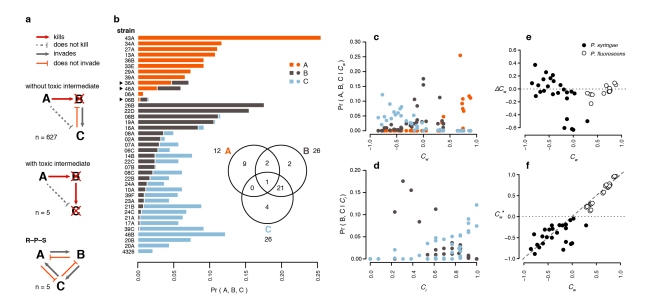


Figure 3: