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### LETTER

# Facilitation promotes invasions in plant-associated microbial communities

#### Abstract

Mei Li,<sup>1,†</sup> Zhong Wei,<sup>1,†</sup>\* Jianing Wang,<sup>1</sup> Alexandre Jousset,<sup>1,2</sup> Ville-Petri Friman,<sup>1,3</sup> Yangchun Xu,<sup>1</sup> Qirong Shen<sup>1</sup>\* and Thomas Pommier<sup>4</sup> While several studies have established a positive correlation between community diversity and invasion resistance, it is less clear how species interactions within resident communities shape this process. Here, we experimentally tested how antagonistic and facilitative pairwise interactions within resident model microbial communities predict invasion by the plant–pathogenic bacterium *Ralstonia solanacearum*. We found that facilitative resident community interactions promoted and antagonistic interactions suppressed invasions both in the lab and in the tomato plant rhizosphere. Crucially, pairwise interactions reliably explained observed invasion outcomes also in multispecies communities, and mechanistically, this was linked to direct inhibition of the invader by antagonistic communities (antibiosis), and to a lesser degree by resource competition between members of the resident community and the invader. Together, our findings suggest that the type and strength of pairwise interactions can reliably predict the outcome of invasions in more complex multispecies communities.

#### Keywords

Antagonism, community ecology, facilitation, invasion resistance, microbe-microbe-plant interactions, rhizosphere.

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#### INTRODUCTION

The characteristics of both resident communities and the invading species are important for determining the outcomes of biological invasions (Williamson & Fitter 1996; Catford et al. 2009). From the resident community perspective, species diversity may be considered a shield to invasions and this effect is often attributed to competition for existing resources (Fridley et al. 2007; Theoharides & Dukes 2007; van Elsas et al. 2012; Wei et al. 2015) where highly diverse communities are thought to efficiently use all the available resource niches leaving no free space for invaders (Case 1990; Tilman 2004). In reality, diversity-invasion resistance relationships are more varied ranging from having neutral to even negative effects (Shea & Chesson 2002; Mallon et al. 2015a; Mehrabi et al. 2016) and are sensitive to environmental conditions (Davis et al. 2000; Roscher et al. 2009; Jousset et al. 2011; Mallon et al. 2015b). Furthermore, it has been shown that trophic network architecture (Wei et al. 2015), species identity effects (Yang et al. 2017) and food web connectance (Smith-Ramesh et al. 2017) are important predictors of invasions and are often linked with community diversity. For example, how species interact might be more important than the number of interacting species within the community (Wei et al. 2015), while invasion resistance may be mediated by certain keystone taxa (Yang et al. 2017). However, the type and strength of resident species interactions have often been overlooked in the context of diversity-invasion resistance.

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Resident species communities form complex ecological webs where multiple species may interact positively or negatively with each other (Kéfi et al. 2012). Positive interactions between species at the same trophic level can result from facilitation or metabolic cross-feeding, where species benefit from the presence of each other (Mulder et al. 2001). Negative interactions may result from resource competition (Wei et al. 2015) or direct interference competition, where species directly suppress each other via antagonism (Bais et al. 2003; Hierro & Callaway 2003; Thorpe et al. 2009; Hu et al. 2016). These interactions may affect the outcomes of invasions in various ways. First, facilitation and competition are likely to affect the resource availability, and hence the availability of free resource niche space, and the likelihood of invasions (Shea & Chesson 2002; Tilman 2004; Stachowicz & Byrnes 2006; Gioria & Osborne 2014; Mallon et al. 2015). It is predicted that highly competitive resident communities are less prone to invasions if they can efficiently utilise and consume resources that would otherwise be available for invaders (Tilman 2004; Jousset et al. 2011; Mallon et al. 2015). This effect is expected to be especially strong in the resident communities that show a high degree of complementarity and hence compete less strongly with each other compared with the invader. In contrast, facilitative interactions between residents could

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potentially increase the number of resource niches via production of secondary metabolites or public goods that can also be utilised by the invader (Stachowicz 2001: Mallon et al. 2015b: Bulleri et al. 2016). Furthermore, competing species can inhibit each other directly by producing toxic metabolites, such as antibiotics. Depending on the spectrum of their activity, antibiotic compounds could have negative effects on both resident community species and the invader (Bais et al. 2003; Hierro & Callaway 2003; Thorpe et al. 2009; Becker et al. 2012; Hu et al. 2016; Wang et al. 2017b). If the invader is particularly sensitive to toxins produced by the resident community, it is expected that antibiotic-mediated interference competition will constrain invasions. In contrast, if toxins have a disproportionally larger negative effect on the members of the resident community, such interference competition is expected to promote invasions (Thorpe et al. 2009; Stubbendieck et al. 2016). Resident community species interactions could further affect certain community-level properties such as ecological stability (Allesina & Levine 2011), which could have indirect effects on invasions (Ghoul & Mitri 2016).

In the present study, we explored to what extent the type (facilitative vs. antagonistic) and strength of two-species resident community species interactions can predict invasions in complex multispecies bacterial communities. Experiments conducted within one trophic level suggest that pairwise bacterial competitions can predict three-species bacterial competitions with as high as 90% accuracy (Friedman et al. 2017). While predicting competitions in species-rich communities might require additional information about potentially emerging higher-order interactions (Friman et al. 2016; Grilli et al. 2017; Levine et al. 2017), these findings suggest that qualitative information regarding species growth in pairwise co-cultures can be used to predict the competitive outcomes of up to 8-species communities (Friedman et al. 2017). Here we extend this approach beyond competition to concurrently explore the role of antagonistic and facilitative resident community interactions for biological invasions (Bruno et al. 2003; Altieri et al. 2010; Traveset & Richardson 2014). Our study system consisted of six non-pathogenic bacterial species (resident community), which were isolated from the tomato plant rhizosphere, and the invader, the plant-pathogenic Ralstonia solanacearum bacterium. Specifically, we first characterised antagonistic and facilitative pairwise interactions within model resident bacterial communities and then directly tested how these interactions predict invasions in more complex multispecies communities both in vitro and in vivo in the tomato rhizosphere. We found that facilitative and antagonistic pairwise interactions reliably predicted invasions: facilitative resident communities were more prone to invasions, while antagonistic resident communities were invaded much less often. Mechanistically, this was linked to direct inhibition of the invader by antagonistic communities (antibiosis), and to a lesser degree by resource competition between the members of the resident community and the invader. Our results suggest that antagonism is an important determinant of community invasion resistance (Case 1990; Tilman 2004), while facilitation might promote invasions by alleviating antagonistic interactions or by releasing vacant niche space for the invader.

#### MATERIALS AND METHODS

#### Bacterial strains and the assembly of resident communities

We used Ralstonia solanacearum strain QL-Rs1115 tagged with the pYC12-mCherry plasmid (Tan et al. 2016) as an invading pathogen in our experiments. Ralstonia solanacearum is a causal driver of bacterial wilt and capable of infecting various economically important crop species (Jiang et al. 2017). We set up model resident communities using six bacterial strains isolated from the tomato rhizosphere at the same location as the pathogen (Qilin [118 ° 57' E, 32 ° 03' N], Nanjing, China). Resident community species listed in Table S1 (Flavobacterium johnsoniae WR4, Chrvseobacterium daecheongense WR21, Delftia acidovorans WR42, Bacillus amyloliquefaciens T-5, Lysinibacillus sphaericus HR92 and Ralstonia pickettii QL-A6) have previously been shown to provide protection for associated host plants by inhibiting R. solanacearum pathogen growth via resource competition or direct toxin production (Figure S1). The resident community composition (Table S2) was manipulated using biodiversity-invasion resistance framework where we modulated both resident community diversity (species richness) and composition and then directly tested how this affected community invasion resistance (Wei et al. 2015). Invasion outcomes were then explained by interactions (1) within resident communities and (2) between resident community and the invader.

## Determining pairwise interactions between resident community species

To quantify the type (facilitative, neutral or antagonistic), strength and direction of each pairwise interaction between resident species, we compared the growth of each species alone and in the presence of each of the other species in twospecies co-cultures (Foster & Bell 2012). All mono-cultures were inoculated with a starting density of 10<sup>5</sup> cells per ml and the co-cultures were inoculated with half of this starting cell density of each species. Resident species were grown for 48 h in liquid NA medium (glucose 10.0 g  $L^{-1}$ , tryptone 5.0 g  $L^{-1}$ , yeast extract 0.5 g  $L^{-1}$ , beef extract 3.0 g  $L^{-1}$ , pH 7.0) in 48well microtiter plates (ending volume of 700 mL per well) at 30 °C with shaking (170 rpm). Bacterial growth was measured as colony number units (CFU) per mL by serial dilution and plating on NA agar plates after 48 h growth. All strains formed distinct colonies on agar plates and could be identified based on colony morphology (Figure S2).

The type of pairwise interaction between two species (here *i* and *j*) was determined by comparing the sum of endpoint of monoculture productivity (population densities) of *i* (MP<sub>*i*</sub>) and monoculture productivity of *j* (MP<sub>*j*</sub>) with the ending productivity of the two-species co-culture ( $CP_{i+j}$ ). As suggested previously, the density of a species mixture is expected to be exactly the sum of their growth in the monocultures if species do not interact (Foster & Bell 2012). Thus, we expected that the interaction between *i* and *j* would be facilitative if  $CP_{i+j} > MP_i + MP_j$ , antagonistic if  $CP_{i+i} < MP_i + MP_i$  and neutral if  $CP_{i+i} = MP_i + MP_i$ .

In order to characterise directionality of pairwise interactions, we compared the ending productivity of each species (CP<sub>*i*</sub> and CP<sub>*j*</sub>) in two-species co-cultures with their ending productivities in monocultures. We then determined the directionality of interaction facilitative if species *j* had a positive effect on *i* ( $\log_{10}(CP_i/MP_i) > 0$ ), antagonistic if  $\log_{10}(CP_i/MP_i) < 0$  and neutral if  $\log_{10}(CP_i/MP_i) = 0$ . We also calculated the mean intensity of facilitation (MIF) of co-cultures as an average of  $\log_{10}$ -transformed pairwise interactions using the following formula:  $MIF_{ij} = \frac{1}{2}[\log(CP_i/MP_i) + \log(CP_j/MP_j)]$ . The two-species community was defined as facilitative when MIF > 0, antagonistic when MIF < 0 and neutral if MIF = 0.

#### Predicting resident species interactions in multispecies communities

We simply assumed that pairwise interactions would not change in the presence of additional species and then predicted resident species interactions in multispecies communities, using two different indices: by calculating (1) the proportion of facilitative pairwise interactions of all possible pairwise interactions and (2) predicted mean intensity of facilitation (PIF) in a multispecies community. For example, among the total number of all possible pairwise interactions of strains i, j and k, if one of these interactions was facilitative  $(CP_{i+i} > MP_i + MP_i)$ , the proportion of facilitative interactions in this resident community was defined as 1/3. Analogous to MIF, we calculated the predicted intensity of facilitation (PIF) in multispecies co-cultures as the sum of log<sub>10</sub>-transformed interactions divided by the number of all possible pairwise interactions within the given community using the following formula:  $PIF = \frac{1}{C_n^2} \sum_{j=1}^{C_n^2} MIF_{ij}$ , where  $MIF_{ij}$  refers the net intensity of one pairwise interaction between species *i* and *j* in a multispecies community, which has a total of  $C_n^2$  number pairwise interactions. The communities were defined as facilitative when PIF > 0, antagonistic when PIF < 00 and neutral when PIF = 0. PIF thus accounted for both the strength and directionality of all potential pairwise interactions in a multispecies community.

#### Validating resident species interactions in multispecies communities

To verify resident species interactions in multispecies bacterial communities, we used qPCR to determine the ending densities of each resident species in monocultures and in all possible co-cultures (3, 4, 5 and 6 resident species communities). All communities were assembled in triplicate in liquid NA medium with a starting density of 10<sup>5</sup> cells per ml in monocultures and 33, 25, 20 and 16.7% of monoculture densities in 3, 4, 5 and 6 resident species communities, respectively. After 48 h in 48-well microtiter plates at 30 °C with shaking (170 rpm), bacterial DNA was extracted using e.Z.N.A. The bacterial DNA kit (OMEGA bio-tek) following manufacturer's protocol and extracted DNA was stored at -80 °C. Species-specific primers were designed for each resident community member (Table S3, Figure S3) and qPCR analyses were carried out with an Applied Biosystems Step One Plus real-time PCR system using SYBR green I fluorescent dye detection in 20 µL volumes with 10 µL of SYBR Premix Ex Taq (TaKaRa Bio Inc., Japan), 2 µL of template, 0.4 µL Dye I, 0.8 µL of both forward and reverse primers (10 mM each) and 6 µL sterile water. The PCR was performed by initially denaturizing at 95 °C for 30 s, cycling 40 times with a 5-s denaturizing step at 95 °C, using a 34-s elongation/extension step at 60 °C, and ending with melt curve analysis at 95 °C for 15 s, at 60 °C for 1 min, and at 95 °C for 15 s. Each resident species community sample was replicated three times.

The observed mean intensity of facilitation (OIF) was calculated using the observed species proportions in the communities based on qPCR data. Similar to PIF, we first determined to what extent the growth of each species was affected by the presence of other species in a given community (growth in the community vs. growth alone). OIF was then calculated according to the following formula:  $OIF = \frac{1}{n} \sum \log(CP_i/MP_i)$ . Communities were defined as facilitative when OIF > 0, antagonistic when OIF < 0 and neutral if OIF = 0. OIF was calculated only based on *in vitro* data and in the case of MIF, PIF and OIF, antagonism included the effects arising from both resource competition and direct inhibition via toxins.

#### Measuring resource competition and direct antagonism between the invader and resident community species

All bacteria were first grown to high densities ( $OD_{600} \approx 1.0$ ) in liquid NA media overnight at 30 °C with shaking (170 rpm), washed three times in 0.85% NaCl, and adjusted to an optical density of 0.5 at 600 nm (OD<sub>600</sub>) with SpectraMax M5 spectrophotometer (Molecular Devices, Sunnyvale, CA, USA). We then measured the growth of the invader and all six resident community species individually on 48 different single-carbon resources (see Table S4) representative of tomato root exudates (Hu et al. (2016)). When the invader and resident community species both grew on the same resource ( $OD_{600} > 0.05$ ), their niches were considered to overlap regarding that given resource. In contrast, when only one strain grew on a specific resource, the niches were considered not to overlap (Wei et al. 2015). This resource competition index estimated the "apparent" resource competition assuming that interacting species would be competing for the same resources even when presented with multiple different resources.

Direct antagonism between the invader and resident community species was measured using supernatant assays (Hu *et al.* 2016). Briefly, after 24 h of growth in NA media, all bacterial monocultures were filtered to remove living cells (0.22 µm filter) after 20 µL of sterile supernatant from each resident species culture was mixed with 180 µl of an overnight-grown *R. solanacearum* culture ( $OD_{600} = 0.05$ , five-fold dilution in liquid NA). The control treatments were inoculated with 20 µL of sterile-filtered NA media instead of bacterial supernatant. All bacterial cultures were grown for 24 h at 30 °C with shaking (170 rpm) before measuring pathogen inhibition as optical density (OD 600 nm). Antagonism was defined as the percentage of reduction in pathogen growth by the supernatant compared to the control treatment for all possible invader-resident species two-species combinations.

#### Measuring invasion success in multispecies communities

#### Invasion success measured in vitro

All possible multispecies resident communities were assembled in triplicate in liquid NA medium with a starting density of  $10^5$  cells per mL (100, 50, 33, 25, 20 and 16.7% of monoculture densities in 1, 2, 3, 4, 5 and 6 resident species communities respectively). Communities were then subsequently exposed to invasion by mCherry-tagged *R. solanacearum* ( $10^4$ cells per ml) in 96-well plates at 30 °C with shaking (170 rpm). After 48 h, total bacterial densities were measured as optical density (OD 600 nm) and invasion success measured as the relative invader density to total bacterial densities using red mCherry protein fluorescence intensity (RFP; excitation: 587 nm, emission: 610 nm) with SpectraMax M5 spectrophotometer.

#### Invasion success measured in vivo

We used a 50-day-long greenhouse experiment with tomato plants to measure invasion success *in vivo*. The soil was collected from a rice field in Wuxi (Jiangsu Province, China), sieved at 5 mm and homogenised and sterilised with gamma radiation. Surface-sterilised tomato seeds (*Lycopersicon esculentum*, cultivar "*Micro-Tom*") were germinated on water-agar plates for 3 days before sowing into seedling plates containing cobalt-60-sterilised seedling substrate (Huainong, Huaian Soil and Fertilizer Institute, Huaian, China). *Ralstonia solanacearum* invasion was tested in all possible two-species resident communities, and due to practical reasons, in 18 multispecies resident communities that varied in their predicted mean intensities of facilitation (Table S5).

Three replicates were used for each resident community, and one replicate consisted of a seedling plate that contained six germinated tomato plants (at the three-leaf stage of growth when grown on 700 g sterilised soil). Similar replication was also used for positive (only the invader) and negative (no bacteria) controls. After 3 days of growth on seedling plates, plants were inoculated with assembled resident communities using root drenching method at a final concentration of  $10^8$  CFU of bacteria g<sup>-1</sup> soil (Wei *et al.* 2013). Seven days after inoculation of resident communities, R. solanacearum was introduced to the roots of all plants at a final concentration of  $10^7$  CFU of bacteria  $g^{-1}$  soil. Tomato plants were then grown for 40 days in a greenhouse (with natural temperature variation ranging from 25 °C to 35 °C) and watered regularly with sterile water. Seedling plates were rearranged randomly every 2 days and disease progression monitored at every 7 days. Forty days after inoculation of R. solanacearum, rhizosphere soil was collected from one plant per replicate seedling tray and the abundance of the invader determined with quantitative PCR as the abundance of R. solanacearumspecific *fliC* gene copy numbers (Hu et al. 2016).

#### Statistical analyses

To meet assumptions of normality and homogeneity of variance, invader densities measured *in vitro* and *in vivo* were log10-transformed. We first assessed the independent effects of the proportion of facilitative interactions and the mean intensity of facilitation based on pairwise resident community interaction on invasions (pathogen density and disease incidence). The type of interaction between resident community species pairs was included into models as a categorical variable (1 = facilitation; 0 = antagonism). In the case of multispecies communities, invasions were explained by three quantitative indices, the proportion of facilitative interactions within a community, the predicted mean intensity of facilitation (PIF) and the observed mean intensity of facilitation (OIF). All indices were fitted as continuous variables and one separate model was used for each index that explained invader densities *in vitro* and *in vivo* and bacterial wilt disease incidence. Additional linear mixed models were used to test invasions as a function of a) niche overlap between resident community, b) mean pathogen inhibition by the resident community and c) resident community species identity effects. All analyses were conducted with SPSS (V. 22) and R (Computing 1991; Team 2013).

#### RESULTS

## Two-species resident species interactions predict invasions in vitro and in vivo

All species had both negative and positive effects on each other while the magnitude and directionality of these effects varied depending on specific species (Fig. 1a). In particular, B. amyloliquefaciens was very antagonistic to the other resident community species. (Fig. 1a). Furthermore, we found that 9 of the communities showed antagonistic, and 6 facilitative pairwise interactions with each other (Fig. 1b, Table S6). On average, facilitative two-species communities reached higher population densities ( $R^2 = 0.79$ , P < 0.001, Figure S4), while antagonistic two-species communities were more inhibitory towards each other ( $R^2=0.32$ , P=0.029, Fig. S5a). No relationship was found between resident species' resource niche overlap and observed mean intensity of facilitation (Figure S5B), which suggests that facilitation did not arise due to niche complementarity. Together these results suggest that the strength of direct inhibition was more important in explaining the type of pairwise interactions between resident community members compared to resource competition.

To link the type of pairwise interaction with the likelihood of invasions, we compared R. solanacearum invasion success in facilitative and antagonistic two-species resident communities. Compared to positive controls (R. solanacearum-only: red dashed line in Fig. 2a-f), pathogen densities were significantly lower in the presence of resident species both in vitro and in vivo. The intensity of pathogen suppression could be predicted by the type of pairwise interactions between the resident species: pathogen density was significantly higher in facilitative compared to antagonistic communities in vitro  $(F_{1,43} = 16.02, P < 0.001, Fig. 2a; R^2 = 0.49, P < 0.0001,$ Fig. 2b) and *in vivo* ( $F_{1,43} = 24.40$ , P < 0.001, Fig. 2c;  $R^2=0.26$ , P=0.0021, Fig. 2d). In line with these results, the bacterial wilt disease incidence was also higher in facilitative compared to antagonistic resident communities ( $F_{1,43} = 9.03$ , P = 0.004, Fig. 2e;  $R^2 = 0.14$ , P = 0.013, Fig. 2f). Mechanistically, this could be explained by loss of pathogen inhibition as suggested by a negative correlation between the mean intensity of facilitation and direct invader suppression ( $R^2 = 0.45$ , P < 0.0001, Figure S6). Together these results suggest that antagonistic two-species resident communities were more



Figure 1 The type and relative strength of resident species pairwise interactions. (a) Network diagram showing the strength and directionality of all pairwise interactions between resident community species. The thickness of lines represents the strength and green and red colour the facilitative or antagonistic effects between different species. (b) Nine of the fifteen pairwise interactions were on average antagonistic (co-culture density < monoculture density) and six facilitative (co-culture density > monoculture density). Panels show two examples: Left, antagonism between species Ba and Cd; Right, facilitation between species Rp and Cd. \*\*\* denotes for statistical significance at P < 0.001. All error bars denote for  $\pm 1$  SEM.

inhibitory not only towards themselves but also against the invader.

#### Predicting and validating invasions in multispecies communities based on pairwise interactions

Interactions within the resident communities could well explain the invader abundance in vitro ( $R^2$ : 0.45, P < 0.0001) and in vivo ( $R^2$ : 0.28, P < 0.0001), and bacterial wilt disease incidence ( $R^2$ : 0.18, P = 0.0002) in vivo (Table 1). The proportion of facilitative interactions were well explained by the increase in invader density in all tested resident communities in vitro ( $R^2 = 0.35$ , P < 0.0001, Fig. 3a). Similarly, both the density of the invader in the tomato rhizosphere ( $R^2 = 0.22$ , P = 0.0004, Fig. 3b) and bacterial wilt disease incidence  $(R^2 = 0.21, P = 0.0004, Fig. 3c)$  increased significantly with increasing proportion of facilitative interactions within the resident communities. The predicted mean intensity of facilitation explained well the increase in invader density in vitro  $(R^2 = 0.45, P < 0.0001, Fig. 3d)$  and in vivo  $(R^2 = 0.21, P < 0.0001)$ P = 0.0005, Fig. 3e) and correlated positively with bacterial wilt disease incidence ( $R^2 = 0.19$ , P = 0.0193, Fig. 3f). The predicted and observed mean intensities of facilitation correlated positively with each other ( $R^2 = 0.44$ , P < 0.0001, Figure S7), demonstrating that pairwise interactions can be used to predict interactions in multispecies communities. As expected, invader densities also increased with increasing observed mean intensity of facilitation both in vitro  $(R^2 = 0.26, P < 0.0001, Fig. 3g)$  and *in vivo*  $(R^2 = 0.17, P)$ P = 0.0019, Fig. 3h). However, the observed mean intensity of facilitation did not correlate significantly with bacterial wilt disease incidence (Fig. 3i).

The low invasion success observed in antagonistic resident communities could be attributed to high levels of direct inhibition of the invader and/or high resource niche overlap between the invader and resident community members. We found that both direct pathogen inhibition and high resource niche overlap reduced invader densities in vitro and in vivo, while only direct pathogen inhibition significantly reduced the disease incidence (Table 1). Direct pathogen suppression correlated negatively with both predicted and observed mean intensities of facilitation suggesting that antagonistic multispecies communities were more inhibitory to the invader (Figure S8). The species B. amyloliquefaciens and F. johnsoniae had strong negative effects on pathogen densities in vitro and in vivo (Table S7). However, only B. amyloliquefaciens had a significant negative effect on disease incidence, while species C. daecheongense had a slightly positive effect on disease incidence (Table S7). Together these results suggest that pairwise resident community interactions can predict invasions in multispecies communities in vitro and in vivo and that these effects were primarily linked with direct pathogen suppression.

#### DISCUSSION

Here, we studied how resident community interactions are linked with invasions in bacterial plant rhizosphere communities. We found that facilitative two- species communities were invaded more easily both in the laboratory and rhizosphere compared to antagonistic resident communities. Furthermore, we could use the pairwise interactions to predict invasion outcomes in multispecies communities containing up to 6 resident species. Specifically, communities characterised by a high proportion of facilitative pairwise interactions, and high predicted and observed mean intensities of facilitation, were more susceptible to invasions. Mechanistically, this was linked to direct inhibition of the invader by antagonistic communities (antibiosis), and to a lesser degree by resource competition between the members of the resident community and the invader. Together these findings suggest that outcomes of relatively simple pairwise interactions can be used to predict invasions in multispecies microbial communities especially



**Figure 2** The type of pairwise resident community interactions predicts invasions *in vitro* and *in vivo*. (a) The *R. solanacearum* invader abundance in antagonistic and facilitative two-species resident communities measured *in vitro*. (b) The relationship between invader abundance and the mean intensity of facilitation in resident communities measured *in vitro*. (c) The relative invader abundance in antagonistic and facilitative two-species resident communities measured *in vitro*. (d) The relationship between invader abundance and the mean intensity of facilitation in resident communities measured *in vitro*. (d) The relationship between invader abundance and the mean intensity of facilitation in resident communities measured *in vivo* in the tomato rhizosphere. (e) The bacterial wilt disease incidence (%) in antagonistic and facilitation in resident communities 40 days after inoculation of the invader. (f) The relationship between disease incidence and the mean intensity of facilitation in resident communities measured *in vivo* in the tomato rhizosphere. In all panels, the red dashed lines show the baseline for positive control treatments (invader-only). In panels, b, d and f, values below and above zero denote for antagonistic and antagonistic pairwise resident communities, respectively. Two and three stars denote for statistical significance at P < 0.01 and P < 0.001 significance levels, respectively. All the bars denote for  $\pm 1$  SEM.

when antagonism and facilitation are strongly linked with the resistance to invasion.

Invasion resistance has been thus far mainly considered from the perspective of resource competition and niche preemption (Case 1990; Tilman 2004; Theoharides & Dukes 2007; van Elsas *et al.* 2012; Wei *et al.* 2015). Our results suggest that facilitative interactions should also be considered in the context of invasions. While it is difficult to pinpoint the exact mechanism between facilitation and invasion, the most likely explanation is the loss of pathogen inhibition along with the increase in the mean intensity of facilitation (Figure S6). This is in line with a previous finding where the increase in the antagonistic activity was found to increase the invasion resistance of *Pseudomonas* resident communities (Hu *et al.* 2016). Another explanation could be that facilitative resident communities were less efficient at competing for resources with the invader compared to antagonistic resident communities. However, this likely played a relatively small role as resource niche overlap had the only significant negative effect on the invader density when measured *in vitro* and *in vivo* but not on disease incidence (Table 1). It is also possible that our resource competition indices measured *in vitro* overestimated the strength of resource competition or underestimated the size of the niche space in the rhizosphere leading to weak correlation with invasions. Furthermore, facilitative interactions could have increased the niche space in the resident communities in favour of the invader, which could have promoted invasions as a side effect (Bulleri *et al.*)

	Invader relative abundance in vitro			Invader abundance measured <i>in vivo</i>			Disease incidence measured <i>in vivo</i>		
	d.f.	F	Р	d.f.	F	Р	d.f.	F	Р
(a) Interactions within resident communities									
Proportion of facilitative interactions	1	0.02	0.885	<b>1</b>	11.82	0.0009	1	7.01	0.009
Predicted mean intensity of interactions	11	129.8	< 2E-16	<b>1</b>	14.29	0.0003	1	12.66	0.0006
Observed mean intensity of interactions	11	8.18	0.005	<b>1</b>	11.24	0.001	1	1.56	0.215
No. of residuals	167			95			95		
Model summary	$R^2: 0.45$	AIC: -303.69		$R^2: 0.28$	AIC: 111.10		$R^2: 0.18$	AIC: 731.07	
(b) Interaction between community and invader									
Niche breadth									
Niche overlap between the invader and resident communities	↓1	13.76	0.0003	↓1	8.62	0.004	1	1.29	0.258
Direct invader inhibition by resident communities	$\downarrow 1$	79.15	8.881E-16	↓1	5.24	0.024	↓1	12.46	0.0006
No. of residuals	168			96			96		
Model summary	$R^2: 0.36$	AIC: -2	277.97	$R^2: 0.13$	AIC:128.58		$R^2: 0.13$	AIC: 735.84	

**Table 1** Two different general linear mixed models (GLM) comparing the interactions within resident communities (a), and interactions between community and invader (b) on invader abundance *in vitro* and *in vivo* and disease incidence measured *in vivo* 

All response variables were treated as continuous variables. The table shows the most parsimonious models selected based on the AIC information. The up and downwards arrows denote for positive and negative effects on response variables, respectively.

2016). For example, previous studies have demonstrated that bacteria can show diet preference between different dietary glycans, which can prolong species coexistence in co-cultures (Tuncil et al. 2017). Such dietary preference might leave some resources less utilised, providing an opportunity for invasion (Tilman 1999). It has also been shown that the breakdown of polysaccharides can allow coexistence of species that liberate polysaccharide breakdown products (PBPs), which are consumed by other species that are unable to grow on the polysaccharides alone (recipients) (Rakoff-Nahoum et al. 2014). Facilitative interactions could thus potentially favour the invader if it is unable to grow on the primary substrates on its own (Bruno et al. 2003). While it is difficult to validate these hypotheses based on our data, we found that facilitative communities were more productive in general and reached higher total cell densities when cultured together compared to alone (Figure S4). This supports the idea that facilitative resident species were benefitting from the presence of each other (for example via cross-feeding), which could also have benefitted the invader by creating vacant niche space. The carrying capacity of resident communities could thus be an important predictor of biological invasions (Gosso et al. 2012).

In addition to within-resident community interactions, the interactions between resident communities and the invader were also good predictors of invasions, albeit to a lesser extent (Fig. 3 and Table 1). While it remains unclear what exact compounds were produced by the resident communities, previous studies have shown that soil bacteria are capable of producing a wide variety of antimicrobials that often suppress *R. solanacearum* (Hu *et al.* 2016; Wang *et al.* 2017b). For example, the *B. amyloliquefaciens* T-5 strain used in this study has been shown to efficiently suppress *R. solanacearum* both in the laboratory and plant rhizosphere (Wang *et al.* 2017b) and this strain also had the greatest negative effect on the pathogen densities and disease incidence in this study (Figure S1). In addition, the strain *F. johnsoniae* had a negative effect on

pathogen densities both *in vitro* and *in vivo*. Together these results suggest that pathogen suppression via toxins was likely mediated by the presence of these species.

In general, pairwise resident community interactions predicted well the observed invasion outcomes in multispecies communities. (Fig. 3, Table 1). However, no correlation was found between the observed mean intensity of facilitation and bacterial wilt disease incidence (Fig. 3i). This suggests that while in vitro mechanisms (resource competition and antibiosis) can robustly predict invasions in more complex in vivo environments (Wei et al. 2015; Hu et al. 2016), they do not account for all aspects of more complex natural environments. There are many potential explanations for these discrepancies that should be validated in future studies. First, investigating the role of microbe-mediated plant immunity is important as both pathogenic and non-pathogenic bacteria can trigger or suppress plant immunity (Chen et al. 2017; Rautenbach et al. 2017). Furthermore, several bacterial secondary metabolites involved in pathogen suppression also impact plant immunity: for example, 2, 4-diacetylphloroglucinol (DAPG) produced by fluorescent Pseudomonas spp. (Bulai & Venturino 2017) or lipopeptide surfactins produced by Bacillus subtilis (Wang et al. 2017a) have a such dual-function. Second, the rhizosphere bacterial communities we used were rather simple, and hence, predictions based on pairwise species interactions should be tested in more complex multi-trophic communities in the future. Lastly, our predictive indexes only estimated the mean net effects and did not distinguish if both or only one of the species benefitted and vice versa (Foster & Bell 2012). While this approach seems to be a good predictor of invasion outcomes, accounting for the directionality of interactions and potential emerging higher-order interactions (Friman et al. 2016; Grilli et al. 2017; Levine et al. 2017) is likely to improve these predictions.

In conclusion, our results suggest that qualitative information regarding species growth in pairwise co-cultures can be used to predict the outcomes of invasions in multispecies



**Figure 3** The relationship between invader abundance and disease incidence with predicted and observed mean intensities of facilitation within multispecies communities. (a and b) The relationship between invader abundance and the proportion of facilitative interactions in the resident communities measured *in vitro* and *in vivo*, respectively. (c) The relationship between bacterial wilt disease incidence (%) and the proportion of facilitative interactions in the resident communities measured *in vitro* and *in vivo* respectively. (f) The relationship between bacterial wilt disease incidence (%) and the predicted mean intensity of facilitation in the resident communities measured *in vitro* and *in vivo* respectively. (f) The relationship between bacterial wilt disease incidence (%) and the predicted mean intensity of facilitation in the resident communities (g and h) The relationship between invader abundance and the observed mean intensity of facilitation in the resident communities measured *in vitro* and *in vivo* respectively. (i) The relationship between bacterial wilt disease incidence (%) and the predicted mean intensity of facilitation in the resident communities measured *in vitro* and *in vivo* respectively. (ii) The relationship between bacterial wilt disease incidence (%) and the observed mean intensity of facilitation in the resident communities. In all panels, red dashed lines show the baseline of invader densities in control treatments (invader-only). In panels d–i, values below and above zero denote for competitive and antagonistic resident communities, respectively.

communities. Even though our results can be broadly applied across different biological problems, they could offer direct solutions in the context of crop protection. Bacterial pathogens impose an ever-increasing threat for agriculture (Olson & Stenlid 2001; Choudhary & Johri 2009; Nicol *et al.* 2011) and recent evidence suggests that the rhizosphere microbiome plays an essential role in controlling the onset of disease (Berendsen *et al.* 2012; Lozupone *et al.* 2012). Understanding the characteristics that make certain microbiomes more resistant to invasions could potentially allow one to harness beneficial bacterial communities for crop protection. While recent studies have shown that microbial diversity alone may be such important characteristic (Wei *et al.* 2015; Hu *et al.* 2016) we here suggest that highly antagonistic microbial communities might also be efficient at constraining pathogen invasions.

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#### AUTHORSHIP

ML, ZW, YX and QS designed research; ML and ZW performed research and analysed data; ML, ZW, AJ, VF and TP wrote the manuscript; all authors contributed to the final draft.

#### DATA ACCESSIBILITY STATEMENT

Data available from the Dryad Digital Repository: https://doi.org/10.5061/dryad.1tp6101.

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