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Stochastic processes govern invasion success in microbial communities when the invader is phylogenetically close to resident bacteria.

**Short title:** Microbial invasion in naturally enriched NOB guilds

**Author affiliations:**

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Abstract

Despite recent efforts in determining the determinants of invasion in microbial communities, experimental observations across different ecosystems are inconclusive. While relationships between resident community diversity and invasion success are often noted, community diversity says little about community assembly processes. Community assembly processes may provide a more inclusive framework to explain – and potentially prevent or facilitate- invasion. Here, we let replicate nitrite-oxidizing bacterial guilds assemble under different conditions from a natural source community and study their compositional patterns to infer the relative importance of the assembly processes. Then, an invader strain from that same guild was introduced at one of three propagule pressures. We found no significant correlation between community diversity and invasion success. Instead, we observed that the effect of selection on invasion success was surpassed by the effect of drift, as inferred from the substantial influence of propagule pressure on invasion success. This dominance of drift can probably be generalized to other invasion cases with high phylogenetic similarity between invader and resident community members. In these situations, our results suggest that attempting to modulate the invasibility of a community by altering its diversity is futile because stochastic processes determine the invasion outcome. Increasing or reducing propagule pressure is then deemed the most efficient avenue to enhance or limit invasion success.

Introduction

Biological invasions can impact resident communities and ecosystems by facilitating fluctuations in biodiversity and in this way alter community function and productivity. For macro-organisms many factors enabling successful invasions have been identified and considerable scientific effort has been devoted to elucidate the determinants of invasion in microbial communities in order to prevent or promote the establishment of new community members (Mallon et al., 2015a; Amalfitano et al., 2014; De Schryver and Vadstein, 2014).
Competition with resident community members has primarily been suggested to determine invasion success, and strong competition decreases invasion success (Fargione and Tilman, 2005; Mallon et al., 2015b; Emery and Gross, 2007). The level of competition is usually inferred from resident community diversity (Elton, 1958) or from the phylogenetic distance between the invader and resident community members (Darwin, 1859). It is suggested that with small phylogenetic distance between invader and resident community members, resident community members impose strong competition on the invader type because phylogenetic similarity implies ecological similarity (Darwin, 1859), which would reduce invasion success (Procheş et al., 2008; Jiang et al., 2010; Thuiller et al., 2010; Tan et al., 2015). In a similar vein, it has been theoretically (Mallon et al., 2015a) and experimentally (van Elsas et al., 2012; Bonanomi et al., 2014; Dillon et al., 2005) suggested that biologically diverse communities are more resistant towards invasion, as originally proposed by Elton (1958). The most commonly cited reason is that more diverse communities are able to utilize resources more efficiently, thus leaving little resource space for invaders, and have higher probability of hosting a type capable of out-competing an invader. However, when community diversity is examined as a single factor determining invasion success without considering the specific context for interpretation, false conclusions are likely (Shade, 2017), because other community assembly processes (i.e., selection, drift, dispersal or speciation) contributing to resident community diversity are often neglected (Kinnunen et al., 2016).

Competition between invader and resident community members is mainly investigated using synthetically assembled microbial communities (De Roy et al., 2013; van Elsas et al., 2012) with limited similarity to natural communities. Synthetically assembled microbial communities allow testing invasion success at different (controlled) diversity levels as well as carefully chosen phylogenetic distances between community members. However, this approach does not allow testing how all community assembly processes affect invasion because oftentimes only one or two processes (selection and/or drift) govern community assembly when establishing synthetic communities with no history of interaction. It is thus
unclear if resident community diversity and phylogenetic distance between invader and resident community members can serve as general predictors of invasion success, beyond synthetic communities. On the other hand, recent studies have suggested that microbial community assembly is more stochastic (Daleo et al., 2009) than recognized in the studies focusing on competition, and that invasion success would primarily depend on propagule pressure (Acosta et al., 2015; Ketola et al., 2017) (the relative abundance of the invader to the resident community), as postulated for communities of macro-organisms (Lockwood et al., 2005; Simberloff, 2009; Von Holle and Simberloff, 2005). While resident community diversity has predicted invasion in several cases (van Elsas et al., 2012; Ketola et al., 2017; Dillon et al., 2005), a similar amount of evidence supports propagule pressure as determinant of invasion (Ketola et al., 2017; Acosta et al., 2015). The lack of consensus across studies may be because the investigations are often limited to only one determinant of invasion. For example, sometimes communities with different diversities are subject to invasion at single propagule pressure (Chapelle et al., 2015; Eisenhauer et al., 2013; van Elsas et al., 2012; Dillon et al., 2005; Jiang et al., 2010), or the phylogenetic distance between resident community members and invader is so large that it is highly improbable that it accurately represents competition for an ecological niche (Bonanomi et al., 2014).

Hence, here we subject guilds of nitrite-oxidizing bacteria (NOB) to invasion by a NOB strain and thus investigate invasion outcome in communities where competition is expected, and where phylogenetic distance between invader and resident community members is low. We hypothesize that with low phylogenetic distance to the resident community members, invasion success is influenced by propagule pressure. Since low phylogenetic distance can confer ecological similarity, neither the resident community members nor the invader would have a competitive advantage and the effect of drift would govern invasion success.

**Materials and methods**

**Invader cultivation**
A culture of *Nitrotoga* HW29 was used as the invader, grown according to its enrichment conditions (Hüpeden *et al.*, 2016) in 250-mL cell culture flasks over a three-month period. After three months, NOB mineral medium was replaced with sterilized non-chlorinated tap water for one month to adjust the invader to the conditions of the resident community. No changes in nitrite removal dynamics were observed in response to this change in the medium. Before the onset of the invasion, all batch cultures were combined and the cell density was determined using a Thoma cell-counting chamber. Then, dilutions of the culture in tap water were spiked with either 0.3 mM or 0.03 mM nitrite to introduce the invader to resident communities with high and low nitrite loading, respectively.

**Invasion in flow-through microcosms**

The experimental set-up consisted of 40 parallel flow-through microcosms. Biofilms developed on Filtralite NC 0.8-1.6 filter material (Saint-Gobain Byggevarer A/S, Oslo) fed with tap water spiked with nitrite at a constant flow rate of 0.43 L/day under ambient temperatures (23 to 25°C). One set of 20 replicates was fed with tap water with 0.3 mM NO₂⁻-N addition while another set of 20 replicates received 10-fold lower nitrogen concentration, 0.03 mM NO₂⁻-N. Resident community biofilms were allowed to develop for 60 days, after which 4 random columns were destructively sampled and used as before invasion reference (called ‘initial’ in results and discussion) and as inocula for batch microcosms (see below) while the remaining columns were subjected to invasion. Three different propagule pressures were applied; such as the total invader cells after a 24 hours of continuous invasion were estimated to represent on average 1%, 10% and 100% of resident NOB cells. The absolute abundance of the resident NOB cells before invasion was estimated from the nitrite removal dynamics and average NOB growth kinetics according to Rittmann and McCarty (1980). We observed complete nitrite removal from day 30 onwards, resulting in total of 0.4mg NO₂⁻N consumed by the resident bacteria at low nitrite loading, and 4mg NO₂⁻N at high nitrite loading, yielding approximately 10⁷ and 10⁸ cells per microcosm at low and high nitrite loading, respectively. Each propagule pressure treatment consisted of 4 replicates whereas 4 replicates at both nitrogen-loadings were maintained as controls without invader (referred to as ‘none’). The flow-through columns were operated
for another 14 days following the invasion after which all material was harvested (‘Final after invasion’ – or ‘Final’) and DNA extracted.

**Invasion in batch microcosms**

Batch microcosms were established in 250-mL cell-culture flasks with the same nitrite concentrations as in the flow-through microcosms. Nitrite-spiked sterile tap water was used as medium and 0.5 g of wet filter material from the initial community from either high or low nitrite loading flow-through columns was added as inoculum. The flasks were subject to rigorous shaking to detach the cells from the filter material and promote growth in suspension. We assumed the absolute abundance of inoculated resident NOB cells to correspond to the abundance estimated for the flow-through microcosms, but corrected for filter material used for inoculation ($10^3$ cells/ml and $10^4$ cells/ml in low and high nitrite loading batch microcosms, respectively). We used this to determine the correct propagule pressure with similar ratios as in the flow-through microcosms: 1%, 10% and 100% of average resident NOB cells. In batch microcosms, the invader cells were added at the same time as the inoculum filter material. The absolute abundance of resident NOB cells differed in flow-through and batch microcosms. Therefore, our experiments included 6 different conditions of absolute propagule pressure, and 3 of relative propagule pressure.

The nitrite removal was measured regularly and when depleted, half of the medium was replaced. After 5 transfer events, the cells were recovered by filtering the total microcosm volume and the retentate was subjected to DNA extraction.

**DNA extraction**

DNA from was isolated using the FastDNA™ SPIN Kit for Soil and the FastPrep® Instrument (MP Biomedicals, Santa Ana, CA) according to the manufacturer’s instruction at room temperature. DNA from liquid microcosms (batch and invader culture) was isolated after filtering (100 mL of invader cell culture and all 250 mL of the batch microcosms) through sterile 0.1 um filters using DNeasy® PowerWater® kit (QIAGEN, Hilden, Germany) according to manufacturer’s instructions at room temperature. The concentration and
purity of extracted DNA were checked using NanoDrop™ 2000 Spectrophotometer (Thermo Fisher Scientific, Wilmington, DE). DNA was then stored at -20°C for subsequent molecular analyses.

**qPCR**

Real-time qPCR assays were performed with a Roche LightCycler® 96 Instrument (Basil, Switzerland). Reaction mixtures (25 μl) contained 12.5 μl SYBR® Green qPCR Mastermix (iQ™ SYBR® Green Supermix; Bio-Rad, Hercules, CA) 1 μl forward and reverse primers (20 μM), 5 μl of template DNA (adjusted to 2 ngDNA μl⁻¹) and 5.5 μl PCR-grade water. Total bacteria were quantified based on 16S rRNA gene copy numbers using the Eubacterial primer set 1055f-1392r as described in Terada et al., (2010). On average 2.5 copies of 16S rRNA gene was estimated per cell, according to rrnDB (Stoddard et al., 2015), with the assumption that majority of the community belongs to *Gallionellacea* and *Nitrospiraceae*. *Nitrospira* cells were quantified using Nitrospira-specific qPCR with primer set NTS232f (Lim et al., 2008) and Nsr1264r (Dionisi et al., 2002) targeting the 16S rRNA genes. Cell numbers were calculated assuming a single 16S operon per cell (rrnDB).

New primer set Ntoga118F (5’-CTTTCAGCGGAAAAGAAAACGCA) and Ntoga840R (5’-CTAAGGAAGTCTCCTCCC) was developed for this study to target the 16S rRNA gene of *Nitrotoga* cells. The primers were designed based on *Nitrotoga* amplicon sequences retrieved from previous experiment where *Nitrotoga* was enriched from tap water spiked with nitrite (Kinnunen et al., 2017). The designed primers cover 27% of known *Nitrotoga* in the SILVA rRNA database (including the 16S rRNA of *Nitrotoga* HW29) and 100% of the tap water enriched *Nitrotoga* from previous experiment (Kinnunen et al., 2017). These primers target a 175 bp product that was verified by constructing a clone library of 180 clones, all of which were determined to belong to *Nitrotoga* genus. The 35 cycles of amplification at 94°C for 30s; 63°C for 30s; 72°C for 60s was performed. Followed by the melting curve analysis.

**Sequencing and amplicon library**

Extracted DNA from all samples was PCR-amplified using primer set PRK341F (5’-CCTAYGGGRBGCASCAG-3’) and PRK806R (5’-GGACTACNNGGTATCTAA-3’) for 35 cycles, to amplify the V3-V4 hypervariable
regions (Yu et al., 2005). Purified PCR products were sequenced on the Illumina MiSeq platform at the DTU Multi Assay Core Center (Lyngby, DK).

All raw 16S rRNA gene amplicons were processed following the DADA2 (version 1.0.3) pipeline with default settings (Callahan et al., 2016). These sequence variants were classified based on the SILVA prokaryotic reference database version 123. Invader sequence was determined from the 100% similarity to 16S sequence of HW29 found in NCBI database by phylogenetic analysis of all Nitrotoga sequence variants using given reference (Figure S5). All sequences have been submitted to NCBI Sequence Read Archive under accession number SRP116646.

**Statistical analysis**

All statistical tests were performed in R. The relative abundance of invader sequence variant of all NOB as well as the similarity between biological replicates was determined using phyloseq package (McMurdie and Holmes, 2013). Phylogenetic distances and Bray-Curtis distances were calculated and plotted as NMDS using phyloseq package. Phylogenetic diversity was calculated using PhyloMeasures package (Tsirogiannis and Sandel, 2016). The statistical difference of the phylogenetic diversity between treatments was determined using a Wilcoxon signed-rank test, comparing the non-invaded control groups at different nitrite loadings. The absolute cell numbers obtained by qPCR were compared using two-way ANOVA test, with factors corresponding to nitrite loading rate and propagule pressure. Correlations between descriptive indices and invader relative abundance were determined using linear regression model and the significance of the difference in correlation between treatments was determined using also a two-way ANOVA test.

**Results and discussion**

**Ecological processes governing resident community assembly**

We enriched 40 resident communities from a tap water source community in flow-through microcosms subjected to two nitrite loading regimes to support the coexistence of competing NOB genera (Kinnunen et al., 2017). We described the resident community composition after 60 days of operation (further referred
to as initial community) and 14 days after the invasion event (referred to as final community). The flow-
through microcosms are expected to facilitate selection, drift and dispersal. We also inoculated batch
microcosms with the initial community from the flow-through microcosms to establish a set of microcosms
where community assembly processes were simplified by elimination of dispersal. The composition of the
batch microcosms was characterized after the inoculation of the ‘resident’ community together with the
invader, representing the starting community after the inoculation (initial community), and after five
subsequent transfer events (final community). While adding invader simultaneously with the resident
community can be viewed as co-assembly, and not invasion, here, we emphasize that the inoculum
material originating from the flow-through microcosms already had 60-days of co-evolution and therefore
can be considered as resident community, even when introduced at the same time with the invader cells.
Faith’s phylogenetic diversity of the resident NOB guild was significantly lower at high vs low nitrite loading
(Table 1) in the flow-through microcosms (Wilcoxon test p=0.02) but not in batch microcosms (Wilcoxon
test p=0.15). This low phylogenetic diversity in flow-through microcosms corresponded to resident
communities where *Nitrotoga* dominated over *Nitrospira* at high nitrite loading, and was consistent with
known differences in nitrite affinity and specific growth rates of these two genera (Nowka *et al.*, 2015). As
pointed out above, diversity indices without context do not say much regarding the ecological processes
shaping the resident NOB guilds. Therefore, in this study, we elaborated on the relative contribution of the
four processes (i.e. selection, drift, dispersal and speciation) that govern community assembly (Vellend,
2010), and subsequently determine invasion outcome.

In Table 1 we provide an overview of the evaluation of the importance of selection, drift and dispersal in
the resident communities. We can neglect speciation, as it is unlikely in the short timeframe of the
experiment that new types arise and achieve significant abundance. Our interpretation of the strength of
processes acting on the resident communities is based on the dynamics and consistency across replicates of
the composition of the non-invaded control communities (Figure 1, Figure S1 and Figure S2) and a
conceptual synthesis of community ecology (Vellend, 2010). We measure stochastic effects as within-group
distances of replicate communities, such that large dissimilarities represent strong effect of stochastic
community assembly processes. Similarly, small dissimilarities between replicate communities point
towards strong effect of selection, as suggested in Evans et al. (2017).

Dispersal was relevant only in flow-through microcosms since they were open to the environment, in
contrast to the batch microcosms, which were fed sterile tap water spiked with nitrite. Dispersal can
influence the diversity, composition, as well as functioning of a community and the effect of dispersal
seems to be enhanced in smaller communities (Zha et al., 2016). For NOB guilds newly assembled from tap
water the contribution of dispersal is low, compared to the contribution of selection and drift (Kinnunen et
al., 2017). Hence, we focus on the relative importance of selection and drift from here on.

The similarity between the resident communities independently assembled from the same source
community was highest in resident communities assembled under flow-through conditions (Figure 2),
which indicates that selection was most important. The direction of selection was affected by the nitrite
loading, as seen from the difference in the ratio of Nitrotoga to Nitrospira at different nitrite loadings. At
the time of invasion Nitrospira abundance had not reached steady state (Figure 1) since it increased
significantly during the 14 days after the invasion event as seen by comparing the ‘Initial Resident’ and
‘Final’ community fractions (ANOVA low nitrite p=0.01; high nitrite p=0.05). In low nitrite loading flow-
through microcosms, selection pressure was positive towards Nitrospira, as Nitrospira increased in
abundance relative to Nitrotoga. Even though Nitrospira also increased significantly in abundance in high
loading flow-through microcosms, the Nitrotoga-to-Nitrospira ratio was higher than in the low nitrite
loading, indicating strongest selection towards resident Nitrotoga. While one Nitrotoga type has been
found to be one of the key nitrite-oxidizers in wastewater treatment (Lücker et al., 2014), indicating its
adaptability at higher nitrite concentrations, little is known about the nitrite affinity of different Nitrotoga
strains in drinking water communities. Previous studies on competition between Nitrospira and Nitrotoga
in drinking water treatment, however, have also observed the dominance of Nitrospira at low nitrite
loading conditions that is outcompeted by *Nitrotoga* at higher nitrite loading conditions (Albers et al., 2018; Kinnunen et al., 2017). Interestingly, the selection in the batch microcosms favored *Nitrospira* under both loading conditions (see final community on Figure 1 and Figure S2). This can be due to the dynamic nitrite-loading in these microcosms, causing nitrite concentration changes over time, providing niches for NOB with a range of affinities for nitrite. In flow-through microcosms, the nitrite concentration attains steady-state (Figure S1), likely selecting for NOB with a narrower range in substrate affinity. Based on this, we expect the invader *Nitrotoga* strain to be less competitive at low nitrite loading than high nitrite loading, in resident communities dominated by competition.

Next, we estimated the relative contribution of drift to the assembly of the resident communities. In both flow-through and batch microcosms, significantly lower guild abundance was observed at low nitrite than at high nitrite loading, as expected (ANOVA flow-through p<0.0001; batch p=0.04). Communities with low abundance are theoretically more affected by drift than communities with more members (Nemergut et al., 2013). The higher dissimilarities between replicate communities after 60 days of low vs high nitrite loading also support this (Figure 2). The contribution of selection over drift was inferred to be highest in high nitrite loading flow-through microcosms based on the high similarity in composition of communities independently assembled from the same source community (Figure 2). In contrast, the contribution of selection over drift was inferred to be lowest in batch microcosms (Figure 2). In these microcosms, half of the community was regularly removed, promoting higher turnover in replacement of removed community members and amplifying the effect of drift compared to the flow-through system.

We can now explain the underlying causes for the differences in the observed phylogenetic diversity of resident NOB guilds (Table 1) based on the community assembly processes discussed above: we saw no significant difference in NOB phylogenetic diversity in batch microcosms, supporting our interpretation that similar processes dominate the community assembly in batch microcosms irrespective of the nitrite loading
regime. In flow-through microcosms, however, the influence of selection over drift varied between the two nitrite loading treatments: in the high nitrite loading microcosms higher selection to drift ratio resulted in significantly lower phylogenetic diversity (Wilcoxon test p=0.02) and the dominance of few community members with high relative fitness.

Successful establishment of the invader

In the flow-through microcosms The resident NOB guild was subject to continuous invasion during a 24 hour period by a culture of *Nitrotoga* HW29 (Hüpeden et al., 2016) while the invader was introduced simultaneously with the resident community in the batches. In flow-through microcosms, the invader strain was subjected to competition with 2 other *Nitrotoga* and 6 *Nitrospira* types at low loading conditions and 3 *Nitrotoga* and 3 *Nitrospira* types at high loading conditions (Figure S2). We used three defined concentrations of invader cells (see invader qPCR data on Figure 1) to achieve low, medium and high relative propagule pressure conditions (estimated to be equivalent to 1%, 10% and 100% of the total resident NOB population), with the aim to test the effect of drift on invasion success. Following the introduction of the invader strain, we allowed another five biomass turnover times (approximately 14 days, estimated from the resident community cell numbers and nitrite loading rates) before sampling the follow-through microcosms. This time for establishment ensured that, if observed, invader persistence would indicate an active population rather than residual invader cells.

Figure 1 shows that the invader cell addition did not significantly change the total NOB cell numbers after 5 biomass turnover times, except in the low nitrite loading batch microcosms (Wilcoxon test p=0.02). The resident community displayed complete nitrite removal during 30 days before the invasion event (Figure S1), suggesting that the resident community had reached its carrying capacity by the time of the invasion event. Hence, if established, the invader *Nitrotoga* displaced some of the resident NOB types or established at low relative abundance.
Based on amplicon sequencing, we could monitor the establishment of the invader strain – as its sequence was not present in the original resident community. The invader Nitrotoga strain was only established in the flow-through microcosms at high propagule pressure, whereas in batch microcosms, it was established at almost all propagule pressures, although at different relative abundance (Table 1). The frequency of establishment increased in both batch and flow-through microcosms with increasing propagule pressure (Figure 3).

Descriptive indices of community composition fail to predict invasion success

First, we tested the diversity-invasibility hypothesis in NOB guilds. We determined correlations between the relative abundance of invader (relative to total NOB) and the phylogenetic diversity of the resident community (Figure S6) as well as invader relative abundance and nearest (Figure S7) and mean phylogenetic distance to the resident community members (Figure S8). We need to emphasize that comparing the guild diversity and phylogenetic distance between invader and resident community is only appropriate for replicate microcosms assembled by similar processes, because different assembly processes contribute differently to community diversity as well as invasion success. In flow-through microcosms, the communities assembled at high and low nitrite loading were governed by different processes; hence, combining the replicates from different treatments would encourage false conclusions of what governs invasion success. Hence, we determined correlations separately for invasion in communities from high nitrite loading and low nitrite loading flow-through microcosms. Because we inferred no difference in dominating assembly process in batch microcosms, we combined the replicate communities at different nitrite loading regimes. We reject the common hypothesis that invader establishment is negatively correlated with resident community diversity (Figure S6). While we observed a negative trend between the resident community diversity and the relative abundance of the invader after establishment in flow-through microcosms, we failed to observe any significant correlation in flow-through microcosms, contrary to batch microcosms. We observed a significant positive correlation between resident community
diversity and the relative abundance of the invader in batch microcosms. Clearly, phylogenetic diversity of
the resident community is not a universal predictor for invasion resistance in a functional guild.

Another metric used to predict invasion success – the nearest and/or average phylogenetic distance to the
resident community (Gallien et al., 2014) – is assumed to be positively correlated with invasion. However,
because selection acts similarly on community members that are phylogenetically similar (Darwin, 1859),
we hypothesized here that drift would, therefore, determine invasion success when phylogenetic distance
between invader and the resident community is low. We neither saw significant correlation between the
nearest (Figure S7) nor the average phylogenetic distance and relative abundance of the invader (Figure
S8). Our observations indicate that when the relative importance of selection over drift in communities is
low, mean phylogenetic distance to the resident community correlates negatively with invader relative
abundance. Failing to see consistent correlations between invasion success and resident community
diversity as well as phylogenetic distance between invader and resident community members, we
evaluated, for the different microcosms, the prevailing community assembly processes and related them to
the subsequent invasion outcome.

Stochastic processes determine invasion success in NOB guilds

Resident communities, assembled with different dominating processes, were subject to invasion at
different propagule pressures. We did not see a consistent correlation between the average distance from
the resident community and invader relative abundance, hence our observations indicated that selection
did not govern invasion outcome. Although, when drift dominates invasion, incidence and relative
abundance of the invader would increase with propagule pressure. Based on this, we observed support for
drift as governing process of invasion.

First, in batch microcosms, where the selection to drift ratio was lowest, we observed a clear effect of
propagule pressure on invasion outcome (Table 1). Both frequency of invader establishment, as well as
relative abundance of invader increased in response to higher concentrations of added invader cells,
supporting drift as the process governing invasion in batch microcosms.
Second, in flow-through microcosms, successful establishment was only observed at high propagule pressure (Table 1). Failure to establish at lower propagule pressures could be a result of drift supported by the characteristics of a flow-through microcosm, where the actual propagule pressure in the system is lower than the theoretical propagule pressure because invader cells could easily flow through the system without attaching to the biofilm surface. Fewer invader cells are more affected by drift and the probability of extinction is increased, compared to larger populations.

Interestingly, drift explained invasion success also in resident communities where the relative importance of selection was high. This is somewhat unexpected, since in communities governed by selection, the competition caused by the fitness difference between invader and resident community members was expected to govern invasion success. One explanation could be that the high phylogenetic similarity of invader and resident community members reduced competition due to absence of large fitness differences.

Our observations were made using natural communities where phylogenetic distances between community members and the invader are very low compared to many other invasion experiments with synthetic communities where phylogenetic distances are up to 10-fold higher (Naughton et al., 2015; Tan et al., 2015). However, when similarly low phylogenetic diversity and low phylogenetic distance between invader and resident were investigated, a similar conclusion was reached: propagule pressure increased invasion success and phylogenetic diversity had no effect on invasion success (Ketola et al., 2017).

In conclusion, our results suggest that for functional guilds invaded by a guild member, where phylogenetic distance between resident and invader is typically low, stochastic processes govern invasion success, even when the relative importance of selection in the resident community is high. Our results also imply that predicting invasion of functional guilds by a member of the same guild from compositional information is nearly impossible, making futile the precise characterization of the composition of resident communities for this purpose. While regular measurements targeting the relative abundance of possible invader can be used to estimate the probability of establishment, the stochastic nature of drift does not allow predictions with high certainty. These observations need to be verified for functional guilds that have opportunities for
larger ecological differences, to test if we can generalize our findings across any type of invasions in microbial communities.

Supplementary information is available at ISME Journal’s website.

References


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Author contributions

M.K, A.D, and B.F.S designed the experiments. M.K performed the experiments and carried out all molecular analyses; the data analysis was performed by M.K supported by A.D. All co-authors assisted in interpreting the results; M.K initiated the manuscript writing, which was finalized with contributions from A.D, H-J.A and B.F.S.

Competing interests

The authors declare no competing financial interests.
Table 1 – Effect of community assembly processes and propagule pressure on frequency of invader establishment

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<th>Mean phylogenetic distance from invader³</th>
<th>Nearest phylogenetic distance from invader³</th>
<th>Relative propagule pressure</th>
<th>Invader establishment frequency⁴</th>
<th>Rel. abundance of invader (%)⁵</th>
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1Strength of the processes is rated with 0 for no effect to ++++ for highest effect inferred from the community composition and the dissimilarity between biological replicates
2Phylogenetic diversity of the NOB guild calculated using the PhyloMeasures package
3Phylogenetic Mean Pairwise Distance
4Mean Nearest Taxon Distance
5Invader establishment frequency detected in four replicates
6Relative abundance of the invader sequence variant out of total NOB sequence variants ± standard deviation within four replicate communities
Figure 1 – Box-and-whisker plot representing the density of NOB in (A) flow-through and (B) batch microcosms before (initial) and after invasion (final) determined by targeted qPCR. The initial community composition was measured before invasion for flow-through microcosms and after first transfer for batch microcosms. Propagule pressure none refers to the non-invaded control microcosms operated in parallel with invaded microcosms.
Figure 2 – Similarities between non-invaded resident communities independently assembled from the same source community in flow-through and batch microcosms under two nitrite loading, based on nonmetric multidimensional scaling ordinations of Bray-Curtis distances across community structures inferred from total community 16S rRNA amplicon libraries.
Figure 3 – Frequency of invader establishment out of 4 replicate microcosms at different propagule pressure (ratio of invader to resident NOB) and in different experimental microcosms (circles for batch microcosms and triangles for flow-through microcosms)