Dolosigranulum pigrum cooperation and competition in human nasal microbiota

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Abstract

**Background:** Multiple epidemiological studies identify *Dolosigranulum pigrum* as a candidate beneficial bacterium based on its positive association with health, including negative associations with nasal/nasopharyngeal colonization by the pathogenic species *Staphylococcus aureus* and *Streptococcus pneumoniae*.

**Results:** Using a multipronged approach to gain new insights into *D. pigrum* function, we observed phenotypic interactions and predictions of genomic capacity that support a role for microbe-microbe interactions involving *D. pigrum* in shaping the composition of human nasal microbiota. We identified *in vivo* community-level and *in vitro* phenotypic cooperation by specific nasal *Corynebacterium* species. Also, *D. pigrum* inhibited *S. aureus* growth *in vitro*. Whereas, robust inhibition of *S. pneumoniae* required both *D. pigrum* and a nasal *Corynebacterium* together, and not either alone. *D. pigrum* L-lactic-acid production was insufficient to account for these inhibitions. Genomic analysis of 11 strains revealed that *D. pigrum* has a small genome (average 1.86 Mb) and multiple predicted auxotrophies consistent with *D. pigrum* relying on its human host and cocolonizing bacteria for key nutrients. Further, the accessory genome of *D. pigrum*
encoded a diverse repertoire of biosynthetic gene clusters, some of which may have a role in microbe-microbe interactions.

**Conclusions:** These new insights into *D. pigrum*’s functions advance the field from compositional analysis to genomic and phenotypic experimentation on a potentially beneficial bacterial resident of the human upper respiratory tract and lay the foundation for future animal and clinical experiments.

**Keywords:** *Dolosigranulum pigrum, Corynebacterium, Staphylococcus aureus,* *Streptococcus pneumoniae,* microbe-microbe interactions, interspecies interactions, upper respiratory tract, nasal, microbiota, comparative genomics
Background

Colonization of the human nasal passages by *Staphylococcus aureus* or *Streptococcus pneumoniae* is a major risk factor for infection by the colonizing bacterium at a distant body site [1-5]. Interventions that reduce the prevalence of colonization also reduce the risk of infection and transmission, e.g., as in [6, 7]. *S. aureus* and *S. pneumoniae* are major human pathogens that cause significant morbidity and mortality worldwide [8-11]. There are also concerns regarding rising rates of antimicrobial resistance [12] and the potential for long-term effects of antibiotics early in life [13]. Thus, efforts have recently focused on the identification of candidate bacteria that confer colonization resistance against *S. aureus* [14-21] and *S. pneumoniae* [22-25], with particular urgency for *S. aureus* in the absence of an effective vaccine.

*Dolosigranulum pigrum* has emerged in multiple studies of the human upper respiratory tract microbiota, colonizing with or without *Corynebacterium* species, as potentially beneficial and/or protective against colonization by *S. aureus* and *S. pneumoniae* [26-50] (reviewed in [14, 51-54]). Little is known about this Gram-positive, catalase-negative, Firmicute bacterium, first described in 1993 [55]. Microbiota studies sampling either nostrils or nasopharynx show very similar results; therefore, for simplicity, we use nasal or nasal passages to denote the area inclusive of the nostrils through the nasopharynx. *D. pigrum* and *S. aureus* are inversely correlated in adult nasal microbiota [31, 41, 56]. Whereas, in pediatric nasal microbiota, *D. pigrum* and members of the genus *Corynebacterium* are overrepresented when *S. pneumoniae* is absent [26, 33, 40]. Moreover, children with *D. pigrum* colonization of the nasal passages are less likely to have acute otitis media [27, 40] and it has been speculated that *D. pigrum*-dominated
microbiota profiles might be more resistant to invasive pneumococcal disease [45].

Furthermore, *D. pigrum* abundance in the nasal passages is inversely associated with wheezing and respiratory tract infections in infants [28] and abundance of *D. pigrum* with *Corynebacterium* in adults provides greater community stability in the face of pneumococcal exposure [50]. The intriguing inference from these studies that *D. pigrum* plays a beneficial role in human nasal microbiota deserves further investigation.

In contrast to the above, there are very few reports of *D. pigrum* in association with human disease [57-61]. Its frequent identification in human nasal microbiota [26, 30-32, 34-37, 39-45, 47, 48, 62-73] coupled with its rare association with infection are consistent with *D. pigrum* functioning as a commensal, and possibly as a mutualist, of humans—characteristics that support its potential for future use as a therapeutic. However, its metabolism and its interplay with other nasal bacteria remain uncharted territory. Using a multipronged approach, we have made significant advances in these areas. First, we identified specific species of candidate bacterial interactors with *D. pigrum* by analyzing nasal microbiota datasets from adults and children. Second, we used *in vitro* phenotypic assays to show that *D. pigrum* exhibits distinct interaction phenotypes with nasal *Corynebacterium* species, *S. aureus* and *S. pneumoniae*. Third, based on the genomes of 11 distinct *D. pigrum* strains, we identify key predicted functions and auxotrophies in its core genome plus a diversity of predicted biosynthetic gene clusters in its accessory genome. This critical shift to phenotypic and genomic experimentation marks a significant advance in understanding *D. pigrum*, a potential beneficially member of human nasal microbiota.
**Results**

**Individual bacterial species are associated with* D. pigrum* in the nasal microbiota of both adults and children.** *D. pigrum* is the only member of its genus and multiple genus-level 16S rRNA gene-based nasal microbiota studies have identified associations between *Dolosigranulum* and other nasal-associated genera, such as *Corynebacterium*, e.g., [28, 29, 36, 38, 40, 43, 73, 74]. In most cases, the taxonomic resolution in the aforementioned studies was limited to the genus or higher taxonomic levels. Thus, we sought to achieve finer taxonomic resolution and to determine what species are associated with *D. pigrum*. We identified two nostril datasets with V1-V2/V1-V3 16S rRNA gene sequences. These regions contain sufficient information for species-level taxonomic assignment to short-read 16S rRNA gene sequences from most nasal-associated bacteria [26, 41]. After parsing sequences into species-level phylotypes, we interrogated each dataset using Analysis of Composition of Microbiomes (ANCOM) [75] to identify bacterial species that display differential relative abundance in the absence or presence of *D. pigrum* sequences (**Figure 1** and **Table S1**). ANCOM is a commonly used approach for identifying associations that accounts for the compositional nature of sequencing data [75]. In the nostrils of 99 children ages 6 and 78 months [26], *Corynebacterium pseudodiphtheriticum* exhibited increased differential relative abundance in the presence of *D. pigrum*, i.e., was positively associated with the presence of *D. pigrum*, as was *Moraxella nonliquefaciens** (**Figure 1A**). In the nostrils of 210 adults from the Human Microbiome Project (HMP), three *Corynebacterium* species—*C. accolens*, *C. propinquum*, *C. pseudodiphtheriticum*—and an unresolved supraspecies of *C. accolens-macginleyi-tuberculostearicum* were positively associated.
with *D. pigrum* (Figure 1B, panels ii-v). Whereas, *S. aureus* was negatively associated with *D. pigrum* (Figure 1B, panel vi). Our previous analysis of these adult data show these *Corynebacterium* species are the most common *Corynebacterium* species in adult nostrils [41]. Also, all of these *Corynebacterium species* and *D. pigrum* are negatively associated with *S. aureus* in this cohort [41]. Such associations in compositional microbiota data lead to testable hypotheses about possible direct microbe-microbe interactions.

We chose to focus on testing hypotheses about direct interactions between *D. pigrum* and the specific nasal *Corynebacterium* species, as well as between *D. pigrum* and *S. aureus*, for several reasons. First, results from both children and adults identified a positive relationship between *D. pigrum* and individual species of *Corynebacterium* in human nasal microbiota, with a positive association of *D. pigrum* and *C. pseudodiphtheriticum* across age groups (Figure 1 and Table S1). Second, associations between *D. pigrum* and the genus *Corynebacterium* are reported in multiple other human nasal microbiota data sets [28, 29, 36, 38, 40, 43, 73, 74] and, therefore, are more likely to be generalizable and of greater impact for the field. Finally, with respect to possible interactions between *D. pigrum* and *S. aureus*, there is a need to identify potential mechanisms of colonization resistance to *S. aureus* given the lack of an effective vaccine. We then used *in vitro* phenotypic assays to test our hypotheses about direct microbe-microbe interactions.

**Nasal Corynebacterium species can enhance the growth of D. pigrum in vitro.** We hypothesized that the strong positive association between *D. pigrum* and the nasal-associated *Corynebacterium* species might be due to these *Corynebacterium* species
releasing metabolites that enhance the growth of *D. pigrum*. As a crude test of this, we quantified *D. pigrum* growth yields on unconditioned agar medium compared to on cell-free agar medium conditioned by growth of *C. pseudodiphtheriticum, C. propinquum* or *C. accolens* (Figure 2). Conditioning agar medium by prior growth of any of these three nasal *Corynebacterium* species increased the yield (measured as colony forming units, CFUs) of two *D. pigrum* strains (CDC 4709-98 and KPL1914) by one to two orders of magnitude compared to growth on unconditioned agar medium (Figures 2A and 2B). Additionally, one strain of *C. pseudodiphtheriticum* (Figure 2A) and the *C. accolens* strain (Figure 2B) increased the growth yield of *D. pigrum* CDC 2949-98, a strain with a higher baseline growth yield. The increases in *D. pigrum* growth yield on the *Corynebacterium* cell-free conditioned agar medium could result from either increased growth rate and/or increased viability, and could be consistent with the nasal *Corynebacterium* species either removing a toxin from the medium or releasing a metabolite that enhances growth and/or survival of *D. pigrum*.

In contrast to the increase in *D. pigrum* growth yield on *C. pseudodiphtheriticum* cell-free conditioned agar medium (Figure 2A), there was no increase in *C. pseudodiphtheriticum* strain KPL1989 growth yield on *D. pigrum* cell-free conditioned agar medium (Figure 2C). Thus, this growth enhancement goes in one direction from nasal *Corynebacterium* species to *D. pigrum*. This is consistent with unilateral cooperation of nasal *Corynebacterium* species—*C. pseudodiphtheriticum, C. propinquum* or *C. accolens*—with *D. pigrum* in the nostril microbiota and support the observed positive *in vivo* community-level relationships.
The positive association between \textit{C. accolens} and \textit{D. pigrum} in adult nostril microbiota datasets indicates that \textit{in vivo} positive interactions between \textit{C. accolens} and \textit{D. pigrum} prevail (Figure 1B, panel ii). However, \textit{in vitro}, we observed either a positive or a negative interaction between \textit{C. accolens} and \textit{D. pigrum} depending on the assay conditions. Unlike \textit{C. propinquum} and \textit{C. pseudodiphtheriticum}, \textit{C. accolens} is a fatty-acid auxotroph and triolein, a model host epithelial-surface triacylglycerol, serves as a source of needed oleic acid in our assays. We observed increased \textit{D. pigrum} growth yield on a semi-permeable membrane atop \textit{C. accolens} cell-free conditioned agar medium (CFCAM) of Brain Heart Infusion (BHI) supplemented with triolein (BHIT) (Figure 2B). In contrast, \textit{D. pigrum} was inhibited when inoculated directly onto this same \textit{C. accolens} cell-free conditioned agar medium (Table 1). This inhibition is reminiscent of our previous finding that \textit{in vitro} the \textit{C. accolens} triacylglycerol lipase, LipS1, hydrolyzes triacylglycerols releasing free fatty acids that inhibit \textit{S. pneumoniae} [33]. Both \textit{D. pigrum} and \textit{S. pneumoniae} belong to the order Lactobacillales and, based on the closeness of their phylogenetic relationship, we hypothesized that \textit{D. pigrum} might be similarly susceptible to free fatty acids such as the oleic acid that \textit{C. accolens} releases from triolein. Indeed, we observed that oleic acid inhibited \textit{D. pigrum} when we challenged \textit{D. pigrum} with oleic acid using a disk diffusion assay (Table 2). We also challenged \textit{D. pigrum} with varying concentrations of oleic acid spread onto plates of BHI agar medium. Similar to the membrane-mediated effect in the \textit{C. accolens} CFCAM experiment above, we observed \textit{D. pigrum} growth at higher concentrations of oleic acid when inoculated onto a semi-permeable membrane atop the oleic-acid-coated medium versus inoculated directly onto the oleic-acid-coated medium (Table 3). This indicates
the membrane provides some protection from inhibition by oleic acid. Overall, these in vitro data indicate that *C. accolens* can both inhibit the growth of *D. pigrum* by releasing antibacterial free fatty acids from host triacylglycerols, such as oleic acid from triolein, (Tables 1 and 2) and enhance the growth of *D. pigrum* by releasing an as-yet unidentified factor(s) (Figure 2B). Collectively, these results point to a complex set of the molecular interactions between these two species.

**D. pigrum inhibits S. aureus growth.** In the absence of a vaccine against *S. aureus*, there are multiple ongoing efforts to identify commensal bacteria that provide colonization resistance to *S. aureus* [15-21, 56] (reviewed in [14]). The ANCOM analysis of the adult nostril microbiota dataset revealed a negative association between *S. aureus* and *D. pigrum* (Figure 1B, panel vi), consistent with previous work [31, 41, 56]. Direct antagonism would be the simplest mechanism underpinning this observation. Therefore, we assayed for the effect of 10 different strains of *D. pigrum* on *S. aureus*. We gave *D. pigrum* a head-start to compensate for its slower growth rate in vitro. *S. aureus* growth was inhibited when it was inoculated adjacent to a pregrown inoculum of each of these 10 *D. pigrum* strains on agar medium (Figure 3).

**D. pigrum production of lactic acid is unlikely to be the primary mechanism for negative associations with S. pneumoniae or S. aureus.** *D. pigrum* lactic acid production has been proposed as a mechanism to explain epidemiologic observations of negative associations between *D. pigrum* and *S. pneumoniae* [74]. Under nutrient rich conditions in vitro, three tested strains of *D. pigrum* produced from 5.7 to 8.2 mM of L-lactic acid with strain KPL1914 producing the highest concentration (Figure 4A). Therefore, we assayed for growth of *S. pneumoniae* in *D. pigrum* KPL1914 cell-free
conditioned medium (CFCM) and in BHI broth supplemented with varying concentrations of L-lactic acid. Three of the four *S. pneumoniae* strains tested showed some growth in 22 mM lactic acid (Figure 4B), and all strains displayed more growth in BHI supplemented with 11 mM L-lactic acid than in the *D. pigrum* KPL1914 CFCM, which had 7.5 mM of *D. pigrum*-produced L-lactic acid (Figure 4B). Thus, the restriction of *S. pneumoniae* growth in *D. pigrum* CFCM is unlikely to be due to *D. pigrum* production of lactic acid. More likely, it reflects competition for nutrients since fresh medium was not added to the CFCM, which, therefore, would have a lower concentration of sugars than BHI broth. However, *D. pigrum* production of a toxin and/or an antipneumococcal compound in BHI broth cannot be excluded. These results indicate that *D. pigrum* production of lactic acid in human nasal passages is unlikely to be the primary molecular mechanism underlying the decreased relative abundance of *S. pneumoniae* in children’s nasal passages when *D. pigrum* is present.

*D. pigrum* is negatively associated with *S. aureus* in adult nostrils [31, 41, 56] and *D. pigrum* excreted a diffusible activity that inhibited *S. aureus* growth on BHI agar (Figure 3). Therefore, we also tested the *in vitro* effect of L-lactic acid on two strains of *S. aureus*. Both showed some growth in 33 mM lactic acid (Figure 4C). Thus, under the tested conditions *D. pigrum* does not produce enough L-lactic acid to restrict *S. aureus* growth. In contrast to *S. pneumoniae*, we would not expect depletion of sugars to have a large effect on *S. aureus* growth in *D. pigrum* CFCM given its broader repertoire of energy source utilization options, e.g., amino acids, and indeed both *S. aureus* strains showed little decrease in growth in *D. pigrum* CFCM. This also revealed to a difference in *D. pigrum* production of the anti-*S. aureus* activity during growth on BHI agar medium.
(Figure 3) versus in BHI broth (Figure 4C). Excretion of metabolites may vary during growth in liquid versus on agar medium and the mechanism of the *D. pigrum* anti-*S. aureus* activity is yet-to-be identified.

**D. pigrum** and **C. pseudodiphtheriticum** inhibit *S. pneumoniae* growth together and not alone. Since *C. pseudodiphtheriticum* was positively associated with the presence of *D. pigrum* in both children and adults (Figure 1), we investigated the effect of a mixed *in vitro* population of *D. pigrum* and *C. pseudodiphtheriticum* on *S. pneumoniae* growth. Agar medium conditioned with a coculture of *C. pseudodiphtheriticum* strain KPL1989 and *D. pigrum* strain CDC4709-98 inhibited *S. pneumoniae* growth, whereas agar medium conditioned with a monoculture of either *C. pseudodiphtheriticum* or *D. pigrum* alone did not (Figures 5 and S1). This could be due to cocultivation resulting in either a greater level of nutrient competition than monoculture of either commensal alone or in the production of diffusible compound(s) toxic/inhibitory to *S. pneumoniae* by either, or both, *D. pigrum* and/or *C. pseudodiphtheriticum* when grown together. Along with the *Corynebacterium* species enhancement of *D. pigrum* growth yield (Figure 2) and the *D. pigrum* inhibition of *S. aureus* growth (Figure 3), these data indicate that the negative associations of *D. pigrum* with *S. aureus* and *S. pneumoniae* are likely mediated by different molecular mechanisms.

Collectively, these phenotypic data (Figures 2, 3 and 5) support a role for microbe-microbe interactions in shaping the composition of human nasal microbiota. These also strengthen the case for *D. pigrum* being a beneficial bacterium that can provide colonization resistance against pathobionts. To learn more about the functional capacity
and genomic structure of *D. pigrum*, we next turned to genomic analysis, which
provided insights into some of the epidemiologic and phenotypic observations
presented above.

**The genomes of 11 *D. pigrum* strains reveal a small genome consistent with a
highly host-adapted bacterium.** We analyzed one publicly available genome of *D.
pigrum* (ATCC51524) and sequenced 10 additional strains (Table S2), which were
selected to ensure representation of distinct strains (see Methods). To start, we focused
on basic genomic characteristics. The 11 *D. pigrum* strain genomes had an average
size of 1.86 Mb (median 1.88 Mb) with 1693 predicted coding sequences (CDS; Tables
S2 and S3). Approximately 1200 CDS were core (Figures S2 and S3; Table S4) and
exhibited a high degree of nucleotide and amino acid sequence conservation (Figure
S4). In Supplemental Text (section I), we further analyzed synteny of two closed
genomes (Figure S5), did BLAST ring comparisons (Figure S6) and constructed a
core-gene-based phylogeny (Figure S7). The 1.86 Mb genome size is consistent with
*D. pigrum* being a highly host-adapted bacterium with reduced biosynthetic capacities,
which are detailed below and in the Supplemental Text (section II) [76].

**D. pigrum** is a predicted auxotroph for amino acids, polyamines and enzymatic
cofactors. The nasal environment is low and/or lacking in key nutrients such as
methionine [77] and *D. pigrum*’s small genome size is consistent with reduced
biosynthetic capacity. To gain insight into how *D. pigrum* functions in the nasal
environment, we examined all 11 genomes finding evidence of auxotrophy for some
amino acids (e.g., methionine), polyamines (e.g., putrescine and spermidine) and
enzymatic cofactors (e.g., biotin) across all strains. In turn, we identified putative
degradation pathways (e.g., methionine), transporters (e.g., polyamines and biotin) and salvage pathways (e.g., folate) suggesting *D. pigrum* acquires some required nutrients exogenously. The Supplemental Text (section II) contains additional details plus predictions on acquisition of metal cofactors. The auxotrophy predictions may be incomplete, since we were unable to grow *D. pigrum* in a chemically defined medium with all 20 amino acids that was putatively replete based on these predictions. Apparent auxotrophy for a number of required nutrients indicates these must be available either from the host or from neighboring microbes in human nasal passages, e.g., possibly from nasal *Corynebacterium* species.

Whole genome sequencing indicates that *D. pigrum* metabolizes carbohydrates via homofermentation to lactic acid. *D. pigrum* produced lactate during *in vitro* cultivation (Figure 4A). Lactic acid bacteria mainly perform either homo- or heterofermentation of carbohydrates [78]. Therefore, we examined the genomic capacity of *D. pigrum* for carbohydrate metabolism (see also Supplemental Text, section III). *D. pigrum* genomes lacked genes required for a complete tricarboxylic acid cycle, which is consistent with fermentation. Moreover, we identified genes encoding a complete glycolytic pathway in all 11 strains that are consistent with homofermentation. All 11 strains harbored a predicted L-lactate-dehydrogenase (EC 1.1.1.27), which catalyzes the reduction of pyruvate to lactate regenerating NAD+ for glycolysis (GAPDH step), consistent with homofermentation to L-lactate as the main product of glycolysis.

The accessory genome of 11 *D. pigrum* strains contains a diversity of biosynthetic gene clusters predicted to encode antibiotics. Lactic acid production alone appears insufficient to account for the negative *in vitro* associations of *D. pigrum*
with *S. aureus* and with *S. pneumoniae* (Figure 4). To delve further into the genetic capacity of *D. pigrum* for possible mechanisms of inhibition, we explored the accessory genome of the 11 sequenced strains. Consistent with a prior report [57], *D. pigrum* appears to be broadly susceptible to antibiotics (Supplement Text, section IV). What emerged in our analysis was a diversity of biosynthetic gene clusters (BGCs) (Table S5 and Figure S8), including a diversity of BGCs predicted to encode candidate antibiotics. Strikingly, although 10 of 10 strains tested displayed inhibition of *S. aureus* growth in vitro (Figure 3), there was no single BGC common to all 10 strains that might encode a compound with antibiotic activity. Based on this, we hypothesize that *D. pigrum* uses a diverse repertoire of BGCs to produce bioactive molecules that play key roles in interspecies interactions with its microbial neighbors, e.g., for niche competition, and potentially with its host. This points to a new direction for future research on the functions that underlie the positive associations of *D. pigrum* in human nasal microbiota with health and highlights the need to develop a system for genetic engineering of *D. pigrum*.

**Discussion**

*D. pigrum* is associated with health in multiple genus-level compositional studies of human URT/nasal passage microbiota. The above species-level genomic and phenotypic experimental data mark a significant advance in the study of *D. pigrum* and set the stage for future research on molecular mechanisms. Further, these phenotypic interactions are consistent with a role for microbe-microbe interactions in shaping the
human nasal microbiota. In nasal passage microbiota datasets, we identified positive associations of *D. pigrum* with specific species of *Corynebacterium* in adults and children and a negative association of *D. pigrum* with *S. aureus* in adults (Figure 1). We observed phenotypic support for these associations during *in vitro* growth. First, unilateral cooperation from three common nasal *Corynebacterium* species enhanced *D. pigrum* growth yields (Figure 2). Second, *D. pigrum* inhibited *S. aureus* (Figure 3). Our genomic analysis revealed auxotrophies consistent with *D. pigrum* reliance on cocolonizing microbes and/or the human host for key nutrients. Genomic analysis also showed an aerotolerant anaerobe that performs homofermentation to lactate. However, *D. pigrum* lactate production (Figure 4A) was insufficient to inhibit either *S. aureus* (Figure 4B) or *S. pneumoniae* (Figure 4C), and is, therefore, not the sole contributor to negative associations with *S. pneumoniae* and *S. aureus in vivo*. Consistent with the multiple reports of a negative association between *D. pigrum*, usually in conjunction with the genus *Corynebacterium*, and *S. pneumoniae*, we observed that cocultivation of *D. pigrum* and *C. pseudodiphtheriticum* produced a diffusible activity that robustly inhibited *S. pneumoniae* (Figures 5 and S1) whereas monoculture of either did not. Finally, we uncovered a surprisingly diverse repertoire of BGCs in 11 *D. pigrum* strains, revealing potential mechanisms for niche competition that were previously unrecognized and opening up a new line of investigation in the field.

The *in vitro* interactions of *D. pigrum* with *S. aureus* and with *S. pneumoniae* support inferences from composition-level microbiota data of competition between *D. pigrum* and each pathobiont. However, these interactions differed *in vitro*. *D. pigrum* alone inhibited *S. aureus* but *D. pigrum* plus *C. pseudodiphtheriticum*, together, robustly
inhibited *S. pneumoniae*. This points to a more complex set of interactions among these specific bacterial members of the human nasal microbiota, which likely exists in the context of a network of both microbe-microbe and microbe-host interactions. To date, mechanisms for only a few such interactions are described. For example, a *C. accolens* triacylglycerol lipase (LipS1) releases antipneumococcal free fatty acids from model host surface triacylglycerols *in vitro* pointing to habitat modification as a possible contributor to *S. pneumoniae* colonization resistance [33].

Multiple mechanisms could result in *D. pigrum* inhibition of *S. aureus* *in vitro* including nutrient competition, excretion of a toxic primary metabolite or of an anti-*S. aureus* secondary metabolite (i.e., an antibiotic). Initial bioassay-guided fractionation approaches failed to identify a mechanism. However, the diverse repertoire of BGCs among the 11 *D. pigrum* strains is intriguing because it includes predicted lanthipeptides and bacteriocins. For example, 4 of the 11 strains harbored putative type II lanthipeptide biosynthetic gene clusters. These clusters are characterized by the presence of the LanM enzyme, containing both dehydration and cyclization domains needed for lanthipeptide biosynthesis [79]. Alignment of these enzymes with the enterococcal cytolysin LanM revealed conserved catalytic residues in both domains [80]. Cleavage of the leader portion of the lanthipeptide is necessary to produce an active compound and the presence of peptidases and transporters within these BGCs suggests these *D. pigrum* strains might secrete an active lanthipeptide, which could play a role in niche competition with other microbes. Additionally, 8 of the 11 *D. pigrum* genomes examined contain putative bacteriocins, or bactericidal proteins and peptides. Intriguingly, the *D. pigrum* strains (CDC4709-98, CDC39-95, KPL1914) exhibiting the strongest inhibition of
*S. aureus* (Figure 3), were the only strains that contained both a lanthipeptide BGC and a bacteriocin, further indicating that *D. pigrum* may employ multiple mechanisms to inhibit *S. aureus* growth, and if both are required for the *in vitro* inhibition might explain the negative results from bioassay guided fractionation.

Mechanisms are coming to light for how other nasal bacteria interact with *S. aureus*. For example, commensal *Corynebacterium* species excrete a to-be-identified substance that inhibits *S. aureus* autoinducing peptides blocking agr quorum sensing (QS) and shifting *S. aureus* shifts towards a commensal phenotype [81]. Also, the to-be-identified mechanism of *C. pseudodiphtheriticum* contact-dependent inhibition of *S. aureus* is mediated through phenol soluble modulins (PSM), the expression of which increases during activation of agr QS [82]. Within broader *Staphylococcus-Corynebacterium* interactions, *C. propinquum* outcompetes coagulase-negative *Staphylococcus* (CoNS), but not *S. aureus*, for iron *in vitro* using the siderophore dehydroxynocardamine, the genes for which are transcribed *in vivo* in human nostrils [83]. Interphylum Actinobacteria-Firmicutes interactions also occur between *Cutibacterium acnes* and *Staphylococcus* species (reviewed in [14]). For example, some strains of *C. acnes* produce an anti-staphylococcal thiopeptide, cutimycin, *in vivo* and the presence of the cutimycin BGC is correlated with microbiota composition at the level of the individual human hair follicle [84]. Of note, Actinobacteria competition with coagulase-negative *Staphylococcus* species could also have network-mediated (indirect) effects on *S. aureus* via the well-known competition among *Staphylococcus* species (reviewed in [85]), which can be mediated by antibiotic production, e.g., [15-17, 19], interference with *S. aureus* agr QS [18, 20, 86, 87] or extracellular protease activity [88], among other
means [14]. Further rounding out the emerging complexity of microbe-microbe interactions in nasal microbiota, multiple strains of *Staphylococcus*, particularly *S. epidermidis*, inhibit the *in vitro* growth of other nasal and skin bacteria, including *D. pigrum*, via to-be-identified mechanisms [16]. The above points to a wealth of opportunity to use human nasal microbiota as a model system to learn how bacteria use competition to shape their community.

Direct cooperation could contribute to the observed positive associations between bacterial species in epidemiological microbiome studies. Conditioning medium with any of the three nasal *Corynebacterium* species positively associated with *D. pigrum in vivo* in human nasal microbiota (Figure 1) enhanced the growth yield of some *D. pigrum* strains (Figure 2). This is possibly by excretion of a limiting nutrient or by removal of a toxic medium component. The genomic predictions of auxotrophy (above and supplemental text) might favor nasal *Corynebacterium* species providing cooperation to *D. pigrum* by excretion of a limiting nutrient. Indeed, mass spectrometry indicates a number of nutrients are limiting in the nose [77].

There were several limitations of our study. First, we analyzed the genomes of 11 strains that were primarily isolated in the setting of disease. It is unclear whether these strains were contaminants or pathogenic contributors [57]. However, *D. pigrum* strains are infrequently associated with disease [58-61, 89-92]. These 11 *D. pigrum* strains encoded only a few potential virulence factors, which is consistent with *D. pigrum* acting primarily as a mutualistic species of humans. Second, the ongoing search for a fully defined chemical medium permissive for *D. pigrum* growth precluded experimental verification of predicted auxotrophies and further investigation of how nasal
Corynebacterium enhance D. pigrum growth yields. Third, the D. pigrum anti-S. aureus factor has eluded purification and identification efforts with standard chemistry approaches and D. pigrum is not yet genetically tractable, limiting genetic approaches to identify it. Fourth, to date, there is no animal model for nasal colonization with D. pigrum and Corynebacterium species, which stymies directly in vivo testing the hypothesis of pathobiont inhibition and points to another area of need within the nasal microbiome field.

Conclusions

In summary, we validated in vivo associations from human bacterial microbiota studies with functional assays that support the hypothesis that D. pigrum is a mutualist with respect to its human host, rather than a purely commensal bacterium. Further, these phenotypic interactions support a role for microbe-microbe interactions in shaping the composition of human nasal microbiota, and, thus, the possibility of developing microbe-targeted interventions to reshape community composition. The next step will be to identify the molecular mechanisms of those interactions and to assess their role in the human host. Such work could establish the premise for future studies to investigate the therapeutic potential of D. pigrum as a topical nasal probiotic for use in patients with recurrent infections with S. pneumoniae, possibly in conjunction with a nasal Corynebacterium species, or S. aureus, in conjunction with established S. aureus decolonization techniques [93].
Methods

Species-level reanalysis of a pediatric nostril microbiota dataset. Laufer et al. analyzed nostril swabs collected from 108 children ages 6 to 78 months [26]. Of these, 44% were culture positive for *S. pneumoniae* and 23% were diagnosed with otitis media. 16S rRNA gene V1-V2 sequences were generated using Roche/454 with primers 27F and 338R. We obtained 184,685 sequences from the authors, of which 94% included sequence matching primer 338R and 1% included sequence matching primer 27F. We performed demultiplexing in QIIME [94] (split_libraries.py) filtering reads for those ≥250 bp in length, quality score ≥30 and with barcode type hamming_8. Then, we eliminated sequences from samples for which there was no metadata (n=108 for metadata) leaving 120,963 sequences on which we performed de novo chimera removal in QIIME (USEARCH 6.1) [95, 96], yielding 120,274 16S rRNA V1-V2 sequences. We then aligned the 120,274 chimera-cleaned reads in QIIME (PyNAST) [97], using eHOMDv15.04 [41] as a reference database, and trimmed the reads using “o-trim-uninformative-columns-from-alignment” and “o-smart-trim” scripts [98]. 116,620 reads (97% of the chimera-cleaned) were recovered after the alignment and trimming steps. After these initial cleaning steps, we retained only the 99 samples with more than 250 reads. We analyzed this dataset of 99 samples with a total of 114,909 reads using MED [98] with minimum substantive abundance of an oligotype (-M) equal to 4 and maximum variation allowed in each node (-V) equal to 6 nt, which equals 1.6% of the 379-nucleotide length of the trimmed alignment. Of the 114,909 sequences, 82.8% (95,164) passed the -M and -V filtering and are represented in the MED output. Oligotypes were assigned taxonomy in R with the dada2::assignTaxonomy() function.
(an implementation of the naïve Bayesian RDP classifier algorithm with a kmer size of 8 and a bootstrap of 100) [99, 100] using the eHOMDv15.1 V1-V3 Training Set (version 1) [41] and a bootstrap of 70. We then collapsed oligotypes within the same species/supraspecies yielding the data shown in Table S6.

**Microbiota community comparison (Figure 1).** The pediatric 16S rRNA gene V1-V2 dataset analyzed at species level here (Table S6), as well as the HMP adult 16S rRNA gene V1-V3 dataset previously analyzed at species level (Table S7 in [41]) were used as input for the ANCOM analysis, including all identified taxa (i.e., we did not remove taxa with low relative abundance). ANCOM (version 1.1.3) was performed using the presence or absence of *D. pigrum*, based on the 16S rRNA gene sequencing data, as group definer. ANCOM default parameters were used (sig = 0.05, tau = 0.02, theta = 0.1, repeated = FALSE (i.e., Kruskal-Wallis test)) except that we performed a correction for multiple comparisons (multcorr = 2), instead of using the default no correction (multcorr = 3) [75]. The Log relative abundance values for the taxa identified as statistically significant (sig = 0.05) are represented in Figure 1 and also available in Table S1.

**Cultivation from frozen stocks.** Bacterial strains (Tables S2 and S7) were cultivated as described here unless stated otherwise. Across the various methods, strains were grown at 37°C with 5% CO2 unless otherwise noted. *D. pigrum* strains were cultivated from frozen stocks on BBL Columbia Colistin-Nalidixic Acid (CNA) agar with 5% sheep blood (BD Diagnostics) for 2 days. *Corynebacterium* species were cultivated from frozen stocks on BHI agar (*C. pseudodiphtheriticum* and *C. propinquum*) or BHI agar supplemented with 1% Tween80 (*C. accolens*) for 1 day. Resuspensions described
below were made by harvesting colonies from agar medium and resuspending in 1X phosphate buffered saline (PBS). Of note, we primarily use agar medium because in our experience *D. pigrum* exhibits more consistent growth on agar medium than in liquid medium. Likewise, growth on a semi-solid surface is likely to better represent growth on nasal surfaces than would growth under the well-mixed conditions of shaking liquid medium.

**Preconditioning growth yield assays (Figure 2).** To assess the growth yield of *D. pigrum* on a polycarbonate membrane atop media conditioned by *Corynebacterium* spp. each. *Corynebacterium* strain was resuspended from growth on agar medium to an optical density at 600 nm (OD$_{600}$) of 0.50 in 1x PBS. Then 100 μL of each resuspension was individually spread onto a 0.2-μm, 47-mm polycarbonate membrane (EMD Millipore, Billerica, MA) atop 20 mL of either BHI agar for *C. pseudodiphtheriticum* and *C. propinquum* or BHI agar supplemented with Triolein (BHIT) (CAS # 122-32-7, Acros) spread atop the agar medium, as previously described [33], for *C. accolens*. After 2 days of growth, membranes with *Corynebacterium* cells were removed, leaving CFCAM. On each plate of CFCAM, we placed a new membrane onto which we spread 100 μL of *D. pigrum* cells that had been resuspended to an OD$_{600}$ of 0.50 in 1x PBS. After 2 days, membranes with *D. pigrum* were removed, placed in 3 mL 1x PBS, and vortexed for 1 min. to resuspend cells. Resuspensions were diluted 1:10 six times, dilutions were inoculated onto BBL CNA agar with 5% sheep blood and colony forming units (CFUs) were enumerated after 2-3 days of growth. To assess the growth yield of *Corynebacterium pseudodiphtheriticum* on a polycarbonate membrane atop media conditioned by *D. pigrum* strains KPL1914 and CDC 4709-98 were grown for 2 days as
described above. *C. pseudodiphtheriticum* KPL1989 growth yield was then measured as described above.

**Growth of *D. pigrum* directly on BHI agar medium supplemented with triolein and conditioned by growth of nasal *Corynebacterium* species (Table 1).** Onto BHI agar supplemented with 200 U/mL of bovine liver catalase (C40-500MG, Sigma) (BHIC), we spread 50 μL of 100 mg/mL of Triolein (BHICT). We then spread 50 μL of a resuspension (OD<sub>600</sub> of 0.50) of each *Corynebacterium* strain onto a 0.2-μm, 47-mm polycarbonate membrane placed atop 10 mL of BHICT agar in a 100-mm-by-15-mm petri dish. After 2 days, we removed each membrane with *Corynebacterium* cells leaving CFCAM. Using a sterile cotton swab, we then spread either a lawn of *D. pigrum* (from cells resuspended to an OD<sub>600</sub> of 0.50 in 1x PBS) or *S. pneumoniae* (taken directly from agar medium) onto the CFCAM. Each lawn then grew for 1-2 days before documenting growth or inhibition of growth with digital photography.

**Oleic acid disc diffusion assay (Table 2).** A lawn of *D. pigrum* or *S. pneumoniae* was spread onto 10 mL of BHIC agar using a sterile cotton swab as described above. Oleic acid (Sigma-Aldrich) was dissolved to a final concentration of 2 mg/mL, 5 mg/mL and 10 mg/mL in ethanol and then we added 10 μL of each to separate, sterile 0.2-μm, 6-mm filter discs (Whatman), with 10 μL of ethanol alone added to a disc as a control. After allowing the solvent to evaporate, filter discs were placed onto the bacterial lawns which were then allowed to grow for 1 day before measuring zones of inhibition and photographing.
Growth of *D. pigrum* directly on versus atop a membrane on oleic-acid-coated agar medium (Table 3). Oleic was dissolved in 100% ethanol to a concentration of 5 mg/ml and then further diluted 10-fold 5 times in ethanol. For each dilution, 100 μL was spread on top of a separate plate of BHI agar medium. Next, 10 μL of *D. pigrum* KPL1914 and CDC4709-98 each resuspended to OD<sub>600</sub> = 0.3 was inoculated both directly on the oleic-acid-coated agar medium and atop of a 0.2-μm, 47-mm polycarbonate membrane (EMD Millipore, Billerica, MA) on the same plate. After 2 days at 37°C, we assessed and photographed the growth. In addition, for each dilution and strain one spot on the membrane was resuspended in PBS to assess CFU counts after serial dilutions and plating on blood agar plates (see above).

*D. pigrum–S. aureus* side-by-side coculture assay (Figure 3). *D. pigrum* cells were harvested with sterile cotton swabs and resuspended in sterile 1x PBS to a minimal OD<sub>600</sub> of 0.3 then 5 μl drops were individually inoculated on BHI agar medium and incubated for 2 days. *S. aureus* JE2 was grown overnight on BBL Columbia CNA agar with 5% sheep blood and resuspended in PBS to an OD<sub>600</sub> of 0.1. Then 5 μl drops of *S. aureus* were inoculated at different distances from the pregrown *D. pigrum*. Inhibition was assessed daily and photographically documented.

Measurement of L-Lactic Acid Concentration (Figure 4A). *D. pigrum* cells were grown from frozen stocks as above. Cells were then harvested with a sterile cotton swab, resuspended to an OD<sub>600</sub> of 0.50 in 1x PBS and inoculated at 1:25 in BHI broth for overnight growth gently shaking (~50-60 rpm) at 37°C under atmospheric conditions. The overnight culture was then inoculated at 1:25 into fresh BHI broth and grown for 24 hrs at 37°C prior to measuring the lactic acid concentration (mmol/L) using a D-lactic acid assay.
acid/L-lactic acid kit per the manufacturer’s instructions (Cat. no. 11112821035, R-
Biopharm AG).

Growth of *S. aureus* and *S. pneumoniae* in *D. pigrum* cell-free conditioned liquid
medium. (CFCM in Figures 4B and 4C) After growth in BHI, as described for L-lactic
cid measurement, *D. pigrum* KPL1914 cells were removed with a 0.22-µM sterile filter
yielding cell-free conditioned medium (CFCM). *S. aureus* strains Newman and JE2 and
*S. pneumoniae* strains TIGR4, DBL5, 603, WU2 were each grown on BBL Columbia
CNA agar with 5% sheep blood for 1 day, harvested with a sterile cotton swab,
resuspended to an OD$_{600}$ of 0.30 in 1x PBS, inoculated at 1:100 into both *D. pigrum*
CFCM and BHI broth and grown for 19-20 hrs at 37°C in shaking (*S. aureus*; 50 rpm) or
static (*S. pneumoniae*) culture under atmospheric conditions. Growth yield was
quantified as OD$_{600}$ absorbance.

Growth of *S. aureus* and *S. pneumoniae* in BHI broth supplemented with L-lactic
acid. (Lactic Acid in Figures 4B and 4C) Strains of *S. aureus* and *S. pneumoniae* were
grown and harvested as described above for inoculation. BHI broth, supplemented with
L-lactic acid (CAS no. 79-33-4; Fisher BioReagents) at varying concentrations from
11mM – 55 mM, was sterilized through a 0.22-µM filter. After inoculating each strain
separately into BHI broth with L-lactic acid, cultures were grown as described above for
growth in CFCM. Growth yield was quantified as OD$_{600}$ absorbance.

Growth assay for *S. pneumoniae* on BHI agar medium conditioned by mono- vs.
coculture of *D. pigrum* and/or *C. pseudodiphtheriticum* (Figures 5 and S1). *D.
pigrum* and *C. pseudodiphtheriticum* strains were grown from freezer stocks as
described above. Cells were harvested with sterile cotton swabs and resuspended in sterile PBS to an OD\textsubscript{600nm} of 0.5. We then spotted 100 µl of 1:1 mixed resuspension on a polycarbonate membrane (see above) on BHI agar medium containing 400U/mL bovine liver catalase. After 2 days of growth, the polycarbonate membrane with \textit{D. pigrum} and/or \textit{C. pseudodiphtheriticum} was removed from each plate leaving CFCAM. \textit{S. pneumoniae} 603 [101] was grown overnight on BBL Columbia CNA agar with 5% sheep blood as described above and, using a sterile cotton swab, a lawn was streaked onto the CFCAM and allowed to grow for 24 hours. Growth/inhibition was assessed daily and photographically recorded. Imaging was difficult due to the transparency of \textit{S. pneumoniae} lawns.

**Selection of strains and preparation of DNA for whole genome sequencing.** \textit{D. pigrum} KPL1914 was isolated from the nostril of a healthy adult (above). In addition, we selected 9 of 27 \textit{D. pigrum} strains from a CDC collection [57] using an \textit{rpoB}-based typing system with a preference for strains isolated from the nasal passages and/or from children (Table S2). Primers Strepto F MOD (AAACTTGGACCAGAAGAAAT) and R MOD (TGTAGCTTATCATCAACCATGTG) were generated \textit{in silico} by mapping primers Strepto F and R [102] to the \textit{rpoB} sequence of \textit{D. pigrum} ATCC 51524 (genome obtained from NCBI; RefSeq: NZ_AGEF00000000.1) with BLAST [103] and manually correcting misalignments in SnapGene viewer 2.8.2 (GSL Biotech, Chicago, IL). PCR were performed using extracted genomic DNA of \textit{D. pigrum}. PCR conditions were as follows: initial denaturation 95°C for 2 minutes, then 30 cycles of denaturation for 30 seconds at 98°C, annealing at 50°C for 30 seconds, elongation 72°C for minutes and a final extension step at 72°C for 10 minutes. PCR products were cleaned using QIAquick
PCR purification kit (Qiagen, Germantown, MD) and sequence determined by Sanger sequencing (Macrogen USA, Boston, MA, USA). In the genomic analysis, we also included the publicly available genome for *D. pigrum* ATCC 51524, which was sequenced by the BROAD institute as part of the HMP (RefSeq NZ_AGEF00000000.1).

*D. pigrum* strains were grown atop membranes for 48 hrs as described above. Cells were harvested with a sterile tip, resuspended in 50 µl of sterile PBS and frozen at -80°C. Genomic DNA was extracted using the Epicentre MasterPure nucleic acid extraction kit (Epicentre, Madison, WI) per the manufacturer’s instructions. We assessed DNA purity using a Nanodrop spectrophotometer (Nanodrop, Wilmington, DE), concentration using Qubit fluorometer (Invitrogen, Carlsbad, CA) and fragment size/quality via agarose gel electrophoresis.

**Whole genome sequencing, read assembly, and annotation (Table S3).** Genomic DNA was sequenced at the Yale Center for Genome Analysis (YCGA), New Haven, CT, on an Illumina MiSeq platform using mated paired-end (2 x 250 bp) technology, assembled using de Bruijn graph algorithms with Velvet [104] with a kmer size of 139 bp and annotated with RAST with FIGfam release 70 [105] and Prokka [106]. In addition, *D. pigrum* strains KPL1914 and CDC#4709-98 [57] were sequenced on a PacBio RS II (Pacific Biosystems, Menlo Park, CA) and sequences were assembled using HGAP version 3.0 [107]. We used an iterative procedure to error correct the PacBio genomes, which involved mapping Illumina reads to the PacBio genomes until there were no differences detected between the Illumina reads and the PacBio assembly [108]. To estimate the degree of assembly errors and missing content that might contribute to the variation in gene content, we compared the Illumina assembly of KPL1914 with the
Illumina-corrected PacBio assembly of KPL1914 to estimate the possible divergence [109]. Within Illumina assemblies, we identified 139 (1566 vs. 1705) predicted coding sequences as determined by RAST annotation absent in the assembly received by PacBio sequencing. Genomes were deposited at NCBI (GenBank: NAJJ00000000, NAQW00000000, NAQX00000000, NAQV00000000, NAQU00000000, NAQT00000000, NAQS00000000, NAQR00000000, NAQQ00000000 and NAQP00000000 in BioProjects PRJNA379818 and PRJNA379966).

Identification of the *D. pigrum* core, shell and cloud genome based on Illumina-sequenced genomes from 11 strains (Figures S2 and S3 and Table S4). Core proteins from RAST-annotated GenBank-files were determined using the intersection of bidirectional best-hits (BDBH), cluster of orthologous (COG) triangles and Markov Cluster Algorithm (OrthoMCL) clustering algorithms using GET_HOMOLOGUES package version 02012019 on Ubuntu-Linux [110] excluding proteins with more than one copy in an input species (as single-copy proteins are safer orthologues, i.e., using flag t-11). GenBank files derived from RAST annotation (see above) were renamed with KPL strain names except for strain ATCC51524. As an initial control, amino acid fasta files (*.faa) were used for the determination of core proteins. We determined the cloud, shell and core genome of each of the 11 sequenced *D. pigrum* strains using the parse_pangenome_matrix.pl script (./parse_pangenome_matrix.pl -m sample_intersection/pangenome_matrix_t0.tab -s) of the GET_HOMOLOGUES package version 30062017 [110]. Definition of cloud, shell and core genome were based on [111]. In brief, cloud is defined as genes only present in a 1 or 2 genomes (cut-off is defined as the class next to the most populated non-core cluster class).
core genome is composed of clusters present in all 11 strains, soft core contains clusters present in 10 genomes and shell includes clusters present in 3 to 9 genomes. Synteny analysis (Figure S5) on BDBH core (with flag t11) was performed using the compare_clusters script (-s) and synteny visualization was done in MAUVE using standard settings [112] after the KPL1914 genome was reverse complemented and both genomes had the origin set at the beginning of dnaA.

Phylogenetic reconstruction, sequence and protein similarities. A monophyletic (clade) core genome phylogenic tree was constructed by including A. otitis (closest neighbor based on the Living Tree Project [113]) an outgroup (Figure S7B). A phylogenic tree without an outgroup was also constructed similarly (Figure S7A). A. otitis ATCC 51267 contigs were downloaded from NCBI (NZ_AGXA0000000.1) and annotated using RAST (see above). Predicted core proteins common to A. otitis and D. pigrum genomes were identified as described above using GET_HOMOLOGUES package. Alignments were done using a loop with Clustal Omega V. 1.2.4 ($ for filename in *.faa; do clustalo -i "$filename" -o clustalo_out/${filename%coral} -v; done) and resulting alignments were concatenated using catfasta2phyml perl script (https://github.com/nylander/catfasta2phyml) $./catfasta2phyml.pl *.faa --verbose > outv.phy. PhyML 3.0 [114] with smart model selection [115] using Akaike information criterion was used for phylogenetic analysis (maximum-likelihood) with 100 regular bootstrap replicates and FigTree (http://tree.bio.ed.ac.uk/software/figtree/) for tree visualization.

BLAST Ring Image Generator (BRIG) was used for visualization of the other sequenced genomes compared to the closed CDC 4709-98 genome (Figure S6) [116].
amino acid and nucleic acid identity (Figure S4) was calculated using GET_HOMOLOGUES package version 30062017 [110]. In brief, a pangenome matrix was generated using the OMCL algorithm (./get_homologues.pl -d dpig_folder -t 0 -M (OMCL)) for homologues identification. Both, ANI and AAI were calculated with all available clusters (t 0). Commands used: Generation of an AA identity matrix: $ ./get_homologues.pl -d “gbk-files” -A -t 0 -M and CDS identity matrix with the command $./get_homologues.pl -d “gbk files” -a ’CDS’ -A -t 0 -M.

Biosynthetic gene clusters and antibiotic resistance genes (Table S5 and Figure S8). AntiSMASH (antibiotics & Secondary Metabolite Analysis SHell) and ClusterFinder [117, 118] were accessed at https://antismash.secondarymetabolites.org/ using default setpoints. Putative antibiotic resistance genes or mutations in genes conferring antibiotic resistance were predicted using Resistance Gene Identifier (RGI) on the Comprehensive Antibiotic Resistance Database (CARD) [119]. Assembly contigs were submitted at RGI (https://card.mcmaster.ca/analyze/rgi) and only perfect and strict hits were allowed. ResFinder version 2.1. (https://cge.cbs.dtu.dk/services/ResFinder/) with 90% threshold for %ID and 60% minimum length [120].

Statistical analyses. R version 3.6.2 was used for statistical analysis and data visualization. The Wilcoxon rank sum test (equivalent to the Mann-Whitney test) was performed using wilcox.test() with paired = FALSE, alternative = “two.sided”.

List of abbreviations. Analysis of Composition of Microbiomes (ANCOM), Colony Forming Units (CFUs), Cell-Free Conditioned Agar Medium (CFCAM), Cell-Free
Conditioned Medium (CFCM), Brain Heart Infusion (BHI), Brain Heart Infusion supplemented with Triolein (BHIT), Biosynthetic Gene Cluster (BGC).

**Declarations**

**Ethics approval and consent to participate.** We isolated *D. pigrum* KPL1914 and *C. pseudodiphtheriticum* KPL1989 from the nostril of an adult as part of a protocol to study the bacterial microbiota of the nostrils of healthy adults that was initially approved by the Harvard Medical School Committee on Human Studies [121], and subsequently approved by the Forsyth Institute Institutional Review Board.

**Consent for publication.** Not applicable

**Availability of data and material.** The authors declare that all data that support the findings of this study are available within the paper (and its supplementary information files), from publicly available repositories, i.e. GenBank, or from the corresponding authors upon reasonable request. All computer code used in this work is either referenced (for published tools) in the methods section and custom-made code (i.e., loop) is given in the methods section. Further details are available from the corresponding authors on reasonable request.

**Competing interests.** The authors declare no competing interests.

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Medicine P3SMP3_155315 (SDB); by the Novartis Foundation for Medical-Biological Research 16B065 (SDB); and by the Promedica Foundation 1449/M (SDB). Funders had no role in the preparation of this manuscript or decision to publish.

**Authors’ contributions.** Conceptualization: SDB, MMP, KPL. Methodology: SDB, SME, MMH. Investigation: SDB, SME, IFE, YK. Interpretation of data: SDB, SME, MMP, IFE, MH, YK, KPL. Visualization: SDB, SME, IFE. Wrote Original Draft: SDB, SME, KPL. Editing and review: SDB, MMP, SME, IFE, KPL. Supervision: SDB, KPL. Funding Acquisition: SDB, MMP, KPL.

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In contrast to when grown on a semi-permeable membrane, *D. pigrum* is inhibited when grown directly on cell-free *C. accolens*-conditioned BHI agar supplemented with triolein as a source of oleic acid.

<table>
<thead>
<tr>
<th>Conditioning Strain</th>
<th><em>S. pneumoniae</em> 603 (6B)</th>
<th><em>D. pigrum</em> CDC 4709-98</th>
<th><em>D. pigrum</em> KPL1914</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>C. accolens</em> KPL1818</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td><em>C. propinquum</em> DSM44285</td>
<td>0</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td><em>C. pseudodiphtheriticum</em> KPL1989</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

*a*0, no growth; +, growth detected, n≥3
Table 2. Oleic acid inhibits *D. pigrum* growth.

<table>
<thead>
<tr>
<th>Oleic Acid (μg/disc)</th>
<th>ZOI (mm)(^a)</th>
<th>S. pneumoniae 603 (6B)</th>
<th>D. pigrum CDC 4709-98</th>
<th>D. pigrum KPL1914</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>10.3 ± 4.7</td>
<td>12.0 ± 2.9</td>
<td>17.0 ± 2.1</td>
<td></td>
</tr>
<tr>
<td>50</td>
<td>22.0 ± 5.4</td>
<td>26.8 ± 4.4</td>
<td>28.4 ± 7.0</td>
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<tr>
<td>100</td>
<td>26.3 ± 6.7</td>
<td>35.8 ± 4.5</td>
<td>39.4 ± 5.0</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\)Mean ZOI ± SD produced in a disc-diffusion assay. ZOIs were measured as the smallest diameter of inhibited growth and measurements include disc diameter (6 mm). Biological replicates (n=4 for *S. pneumoniae*, n=5 for *D. pigrum*) were averaged.
Table 3. A 0.2-µm, 47-mm polycarbonate membrane provides *D. pigrum* with some protection against inhibition by oleic acid *in vitro*.

<table>
<thead>
<tr>
<th>Oleic acid (µg plated)</th>
<th>Growth directly on agar&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Growth on membrane</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td><em>D. pigrum</em> strain</td>
<td></td>
</tr>
<tr>
<td></td>
<td>KPL1914</td>
<td>CDC 4709-98</td>
</tr>
<tr>
<td>500</td>
<td>0</td>
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<td>0</td>
</tr>
<tr>
<td>5</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>0.5</td>
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</tr>
<tr>
<td>0.05</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>0 (BHI)</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>0 (CSBA)</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

<sup>a</sup>0, no growth, +, growth detected, *n*=3
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Figure Legends.

**Figure 1.** Individual nasal *Corynebacterium* species exhibit increased differential relative abundance in the presence of *D. pigrum* in human nostril microbiota. We used ANCOM to compare species/supraspecies-level composition of 16S rRNA gene nostril datasets from (A) 99 children ages 6 and 78 months and (B) 210 adults when *D. pigrum* was either absent (Dpi-) or present (Dpi+) based on 16S rRNA gene sequencing data. Plots show only the taxa identified as statistically significant (sig = 0.05) after correction for multiple testing within ANCOM. The dark bar represents the median; lower and upper hinges correspond to the first and third quartiles. Each grey dot represents the value for a sample, and multiple overlapping dots appear black. Dpi = *Dolosigranulum pigrum*, Cac = *Corynebacterium accolens*, Caa/Cma/Ctu= supraspecies *Corynebacterium accolens_macginleyi_tuberculostearicum*, Cpr = *Corynebacterium propinquum*, Cps = *Corynebacterium pseudodiphtheriticum*, Mno = *Moraxella nonliquefaciens*. Only three species and one supraspecies of *Corynebacterium* out of the larger number of *Corynebacterium* supraspecies/species present in each dataset met the significance threshold. Specifically, in the adult nostril dataset, there were 21 species and 5 supraspecies groupings of *Corynebacterium* in addition to reads of *Corynebacterium* that were non-assigned (NA) at species level. These data are previously published and visible in Table S7 of reference 42. In the pediatric dataset, there were 16 species of *Corynebacterium* in addition to the (NA) at species level *Corynebacterium* reads (see Table S7 of this manuscript). The Log relative abundance numerical data represented in this figure are available in Table S1.
Figure 2. *D. pigrum* growth yields increase on cell-free conditioned agar medium (CFCAM) from nasal *Corynebacterium* species and not vice versa. Growth yield of *D. pigrum* strains CDC 2949-98, CDC 4709-98 and KPL1914 was quantified as the number of CFUs grown on a polycarbonate membrane placed onto (A) cell-free conditioned BHI agar from *C. propinquum* (aqua green) or *C. pseudodiphtheriticum* (dark and light green) or (B) cell-free conditioned BHI-Triolein (BHIT) agar from *C. accolens* (blue) and compared to growth on unconditioned BHI agar (dark grey) or unconditioned BHIT agar (light grey), respectively. Growth yield of *C. pseudodiphtheriticum* KPL1989 on CFCAM from *D. pigrum* strains (orange) compared to unconditioned medium (white) was assessed similarly (C). BHIT was used for growth of *C. accolens* since it is a fatty-acid auxotroph and releases needed oleic acid from triolein. Preconditioning strains were grown on a 0.2-μm, 47-mm polycarbonate membrane for two days to generate CFCAM. After removal, we then placed a new membrane on the CFCAM onto which we spread 100 μL of target bacterial cells that had been resuspended to an OD$_{600}$ of 0.50 in 1x PBS. After 2 days of growth, CFU were enumerated as described in Methods. CFU counts were compared independently for each individual strain (A and B, n=5) or medium (C, n=4) using a Wilcoxon rank sum test with Bonferroni correction for multiple comparisons to the unconditioned medium. Dark bars represent medians, lower and upper hinges correspond to the first and third quartiles and outlier points are displayed individually. *, $p < 0.05$; **, $p < 0.001$

Figure 3. Ten different strains of *D. pigrum* inhibit methicillin-resistant *S. aureus* USA300 strain JE2. Ten pregrown *D. pigrum* isolates produced a diffusible activity that inhibited the growth of *S. aureus* strain JE2 on BHI agar ($n \geq 3$ independent
experiments). Representative images are shown for each strain. *D. pigrum* was resuspended in PBS then a 5 µl drop was placed onto BHI agar and pregrown for 48 hrs. After that, *S. aureus* JE2 was inoculated adjacent to the *D. pigrum*. Inhibition was assessed after 24 and 48 hrs (48 hrs shown here).

**Figure 4.** Lactate production by *D. pigrum* is insufficient to inhibit pathobiont growth. Strains of *S. pneumoniae* and *S. aureus* grew in the presence of higher levels of L-lactic acid than those produced by *D. pigrum in vitro*. (A) The concentration of L-lactic acid (mM) produced by three *D. pigrum* strains was measured after 24 hrs of gentle shaken aerobic growth in BHI broth at 37°C (*n*=5) as compared to the basal concentration of L-lactic acid in BHI alone (none). (B) The average growth (OD₆₀₀) of 4 *S. pneumoniae* strains in *D. pigrum* KPL1914 CFCM or in unconditioned BHI broth supplemented with different concentrations of L-lactic acid measured after 19–20 hrs of static aerobic growth at 37°C (*n*=4). (C) The average growth (OD₆₀₀) of 2 *S. aureus* strains in *D. pigrum* KPL1914 CFCM or in unconditioned BHI broth supplemented with different concentrations of L-lactic acid measured after 19–20 hrs of shaken aerobic growth at 37°C (*n*=4). Average growth of *S. pneumoniae* in CFCM and 11 mM L-lactic acid were analyzed independently for each individual strain using a Wilcoxon rank sum test. Dark bars represent medians, lower and upper hinges correspond to the first and third quartiles and outlier points are displayed individually except in panel A where dots for all individual sample values are represented. *None of the *S. pneumoniae* or *S. aureus* strains displayed growth in 55 mM L-lactate.*

**Figure 5.** *D. pigrum* and *C. pseudodiphtheriticum* grown together but not *D. pigrum* alone inhibit *S. pneumoniae* in an in vitro agar medium-based assay.
Representative images of *S. pneumoniae* 603 growth on (A) BHI alone or on CFCAM from (B) *C. pseudodiphtheriticum* KPL1989, (C) *D. pigrum* KPL1914 or (D) both *D. pigrum* and *C. pseudodiphtheriticum* grown in a mixed inoculum (n=4). To condition the medium, we cultivated *D. pigrum* and/or *C. pseudodiphtheriticum* on a membrane, which was then removed prior to spreading a lawn of *S. pneumoniae*. For monoculture, 100 μL of either *D. pigrum* or *C. pseudodiphtheriticum*, resuspended to an OD$_{600}$=0.50, were inoculated onto the membrane. For mixed coculture, 50 μL of *D. pigrum* (OD$_{600}$=0.50) were mixed with 50 μL of *C. pseudodiphtheriticum* (OD$_{600}$=0.50) to yield a final volume of 100 μL for the inoculum, such that each bacterial species is present in the coculture inoculum at half the amount used for the respective monoculture inoculum. Images were cropped. Black marks indicate edges of where the membrane had been.
Figure A: Lactic acid (mM) in D. pigrum strain in BHI broth.

Figure B: Turbidity (OD_{600}) for S. pneumoniae strain.

Figure C: Turbidity (OD_{600}) for S. aureus strain.

Each figure shows different lactic acid (mM) in various mediums for different strains.
S. pneumoniae growth

A B
BHI

C D
Cps CFCAM

C D
Dpi CFCAM

Cps + Dpi CFCAM