Identification of Nasal Gammaproteobacteria with Potent Activity against Staphylococcus aureus: Novel Insights into the “Noncarrier” State

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ABSTRACT Staphylococcus aureus nasal carriage provides the bacterial reservoir for opportunistic infection. In comparing the nasal microbiomes of culture-defined persistent S. aureus carriers versus noncarriers, we detected S. aureus DNA in all noses, including those with an established history of S. aureus negativity based on culture. Colonization with Gammaproteobacteria, including Klebsiella aerogenes, Citrobacter koseri, Moraxella lincolnii, and select Acinetobacter spp., was associated with S. aureus noncarriage. We next developed physiological competition assays for testing anti-S. aureus activity of isolated nasal species, utilizing medium modeling the nutrient-limited fluid of the nasal mucosa, polarized primary nasal epithelia, and nasal secretions.

IMPORTANCE Nasal carriage of Staphylococcus aureus is a risk factor for infection, but it is not yet understood why some individuals carry nasal S. aureus persistently, intermittently, or seemingly not at all when tested via culture methods. This study compared the nasal microbiomes of established S. aureus carriers and noncarriers, identified species associated with noncarriage, and tested them for anti-S. aureus activity using assays developed to model the nutrient-limited nasal mucosa. We determined that all nostril swabs contain S. aureus DNA, even swabs from hosts considered to be long-term noncarriers. Select members of the Gammaproteobacteria class were more prevalent in noncarrier than carrier nostrils and demonstrated potent activity against multiple strains of S. aureus. The results described here provide a better understanding of how the nasal microbiome controls S. aureus growth and viability and may be useful in the design of improved S. aureus decolonization strategies.

KEYWORDS Staphylococcus aureus, nasal carriage, microbiome, bacterial interference

Staphylococcus aureus infections have been a concern for health care for decades, especially when considering the rise in antibiotic-resistant strains and postoperative S. aureus infections that plague hospital systems worldwide (1, 2). While S. aureus is not generally considered to be a main colonizer of the human gut (3), it resides at...
least transiently in the moist mucosa of the anterior nares (3–6). *S. aureus* nasal carriage (SANC) is typically asymptomatic, but transmission of *S. aureus* from the nasal cavity to other areas of the body renders carriers more prone to developing *S. aureus* infections (7, 8). Nasally carried *S. aureus* is also frequently spread to household members and domestic pets (9–11). On account of this and the finding that even newborns are *S. aureus* colonized (12), *S. aureus* should be considered ubiquitous, and investigations of carriage dynamics warrant high priority.

Previous studies have suggested that SANC is determined by host and environmental factors rather than by individual strain characteristics of the bacterium (13–15). Higher carriage rates are found in infants, toddlers, and the elderly; risk factors for adults include HIV, type II diabetes, atopic dermatitis, autoimmune disease, rheumatoid arthritis, cancer, prolonged hospital stays, and close living quarters such as prisons, nursing homes, and daycare centers (16–20). Smoking has been linked with elevated nasal *S. aureus* load and a decreased capacity for clearing nasal *S. aureus* (21). Conventionally, SANC is categorized according to frequency of *S. aureus* detection by culture methods during repeat sampling over months and years. Nostril swabs of persistent carriers always contain detectable *S. aureus*, and intermittent carriers periodically clear *S. aureus* to undetectable levels, while noncarriers lack detectable *S. aureus* (22, 23). We demonstrated that nasal fluids of human *S. aureus* carriers have less intrinsic anti-*S. aureus* activity than nasal fluids collected from noncarriers (15, 24, 25) even though the basal components of nasal secretions (e.g., amino acids, salts, organic acids, trace elements, sugars) are not variable between donors (26). Host antimicrobial peptide polymorphisms fail to distinguish *S. aureus* carriers from noncarriers, and HNP1 to -3 and various cytokine levels are elevated in carriers, suggesting that these proteins are more the result of SANC than a reason for noncarrier status (13, 15, 21, 27). Collectively, these insights led us to the current question of how the constantly fluctuating nasal microbiota affect SANC status.

The human nasal microbiome consists mainly of Corynebacterium (phylum Actinobacteria), Staphylococcus (phylum Firmicutes), and Cutibacterium (formerly Propionibacterium; phylum Actinobacteria), balanced by lower levels (typically) of various Proteobacteria as well as Dolosigranulum and Streptococcus genera of the phylum Firmicutes. Nasal microbiome dynamics appear to be less determined by host genetics than by environment, with fluctuations influenced by hygiene, climate, diet, prescription drugs, and interpersonal relationships (28, 29). At the species level, the role of the nasal microbiome in promoting or preventing SANC remains mostly unknown. Conflicting reports have emerged regarding the role of *S. epidermidis* (28, 30, 31), *S. lugdunensis*, *S. hominis*, and *S. saprophyticus* have demonstrated promise as anti-*S. aureus* effectors (31, 32), but it remains unclear if they are present in high enough abundance to distinguish SANC from non-SANC individuals. Corynebacterium pseudodiphtheriticum produces a factor(s) with bactericidal activity toward *S. aureus* in *in vitro* agar-based zone of clearance assays (33). However, the relative abundance of this species was found to be inversely related to *S. aureus* in ciliated respiratory epithelia but not in the anterior nares (34). Here, we evaluated the nasal microbiomes of the three SANC subgroups using 16S rRNA amplicon sequencing and also compared noncarrier versus persistent nostrils at the species level using culture methods and shotgun DNA metagenomic sequencing. Notably, certain Proteobacteria were more strongly associated with noncarriers. We developed physiologically relevant assays for testing nasal *S. epidermidis*, *C. pseudodiphtheriticum*, and several nasal strains of Klebsiella, Citrobacter, Acinetobacter, and Moraxella spp. for inhibitory activity against nasal *S. aureus*.

**RESULTS**

Nasal *S. aureus* load correlates with *in vitro* growth of *S. aureus* in nasal secretions. We demonstrated previously that nasal secretions from *S. aureus* carriers support the *ex vivo* growth of *S. aureus* more than secretions from healthy noncarriers (4, 15, 24).
However, host antimicrobial peptides, including HNP-1, HNP-3, and HBD-2, are elevated in *S. aureus* carrier fluids compared to noncarrier fluids (4, 13). This indicates that host expression of known anti-*S. aureus* effectors is more likely the result of *S. aureus* colonization than a reason that some individuals appear not to carry *S. aureus*. To further evaluate the role of secreted factors in modulating nasal *S. aureus* load, we measured the growth of five different *S. aureus* isolates in *S. aureus* carrier secretions collected at various time points when the corresponding nasal *S. aureus* load was known. Figure 1A and B demonstrate the association between nasal *S. aureus* load and growth of *S. aureus* in matched nasal secretions for two donors (D832 in Fig. 1A and D837 in Fig. 1B). These subjects were in good health and were categorized as persistent *S. aureus* carriers by presenting *S. aureus*-positive nasal swabs weekly for over a year. In 17 nasal fluid samples collected from nasal carriers in which a matched nasal *S. aureus* count was available, the average growth of 5 *S. aureus* strains correlated with the nasal load of the host nose (Fig. 1C; Pearson $r = 0.84$, $P < 0.0001$). Collectively, this suggested that nasal secretions contain factors that dictate how much *S. aureus* is detectable by culture methods. Considering that nasal secretions have been demonstrated to be
indistinguishable between *S. aureus* carriers and noncarriers in terms of nutrient and metabolite composition (26), it became evident that nasal microbiome composition might control nasal *S. aureus* survival.

**Genus-level nasal microbial community profiling reveals no appreciable difference between nasal *S. aureus* carriers and noncarriers.** We next sought to identify the abundant species present in the noses of individuals considered to be *S. aureus* noncarriers as detected by culture methods and to characterize which species levels fluctuate when intermittent carriers cleared or nearly cleared *S. aureus*. Nasal swabs were vortexed in broth and spread onto 6 different agars, including both nonselective (plate count, nutrient, Trypticase soy agar [TSA], TSA II/5% blood) and *S. aureus*-selective (mannitol salt agar or CHROMagar *S. aureus*) formulations. Even in demonstrated *S. aureus* noncarriers, it quickly became apparent that non-*Staphylococcus* genera account for a minority of visible colonies when nutrient-rich agars are employed. Out of 60 colonies (identified by full-length 16S sequencing), only 5 non-*Staphylococcus* spp. were identified (*Micrococcus luteus*, *Micrococcus yunnanensis*, *Pantoea septica*, *Klebsiella aerogenes*, and *Enterococcus faecalis*), while *S. aureus* (in carrier noses), *S. epidermidis*, and *S. hominis* accounted for most colonies identified. In an effort to detect more *Proteobacteria* and *Actinobacteria* species in noses with known SANC status, we next performed microbial community profiling of nasal swab DNA. 16S rRNA amplicon sequencing (Illumina MiSeq) was performed on 65 nasal swab DNA extracts collected from 25 individuals over time—34 swabs from noncarrier nostrils (15 individuals), 9 swabs from persistent *S. aureus* carrier nostrils (4 individuals), and 22 (11 *S. aureus*-negative/11 *S. aureus*-positive) swabs from intermittent carrier nostrils (6 individuals). As shown in Fig. 2A, taxonomic profiling at the genus level revealed that all noses are heavily colonized with *Staphylococcus* and *Corynebacterium* regardless of SANC status and that a majority of noses also contain *Acinetobacter*, *Dolosigranulum*, and a mixture of low-abundance genera from the *Actinobacteria*, *Firmicutes*, and *Proteobacteria* phyla. A stacked bar representation of each nostril’s taxonomic profile (*n* = 65) is provided as Fig. S1. Shannon diversity analysis revealed that noncarriers exhibited more diverse taxa than the other groups, although only the difference between noncarriers and intermittent carriers reached statistical significance (Fig. 2B; and *P* = 0.025). Total bacterial density (CFU/swab) ranged widely for all groups (10^2 to 10^6 CFU/swab) and trended higher for intermittent versus noncarriers (Fig. 2C; and *P* = 0.042), suggesting that noncarriers’ increased microbial diversity was not due simply to enhanced bacterial load. There was no difference in the number of sequence reads between groups (not shown). The 65 samples were clustered based on their genus-level taxonomic profiles using the Dirichlet multinomial mixture (DMM) model framework (35), and this analysis revealed the presence of two clusters, a *Staphylococcus*-prevalent cluster and a *Corynebacterium*-prevalent cluster. Each of these contained a mixture of noncarriers, intermittent carriers, and persistent carriers (not shown). Stratifying the intermittent carrier nostril taxa according to SANC status (11 *S. aureus*-negative versus 11 *S. aureus*-positive swabs) also revealed no appreciable differences between the nasal microbial communities (Fig. S2). Taken together, these data corroborate the findings of Piewngam et al. (36), who also reported that genus-level profiling does not provide the resolution required to distinguish SANC from non-SANC individuals.

**Identification of species associated with *S. aureus* culture-negative noses.** We next took a two-pronged approach to identifying nasal microbiome species that might disfavor *S. aureus* colonization. Six subjects, each of whom had submitted regular swabs for at least 3 months and thus presented a known SANC status, self-collected nasal swabs twice weekly. With the first swab, SANC status was confirmed by identification of *S. aureus* colonies on CHROMagar *S. aureus* plates, and in parallel, isolation of non-*Staphylococcus* species was attempted by culturing nasal samples on MacConkey II and Leeds agars, which favor growth of Gram-negative species. The six subjects comprised 3 noncarriers, 2 persistent carriers, and 1 carrier with an ~80% *S. aureus*-positive swab rate over 3 years of monitoring. In general, noncarrier donor swabs yielded tens to hundreds of colonies on MacConkey II and Leeds agars, while SANC donor swabs yielded no CFU on MacConkey II agar and few CFU on Leeds agar. Dozens of colonies
were plucked and cultured overnight in nutrient broth, followed by identification by 16S PCR/Sanger sequencing. The most common isolate from noncarrier noses was *Klebsiella aerogenes*, followed by *Citrobacter koseri* and *Klebsiella oxytoca*. For the Leeds agar CFU cultured from *S. aureus*-positive swabs, nearly all colonies (18 of 21) were identified as *Staphylococcus* (*epidermidis*, *haemolyticus*, and *hominis*), and no *Klebsiella* or *Citrobacter* species were detected.

Subsequent nasal swab collections, occurring 2 to 3 days after cultured swabs, were immediately processed for DNA isolation coupled with a mammalian DNA depletion step. The resulting DNA was fragmented and indexed, and 14 samples were pooled and sequenced (single-lane HiSeq 2 × 150 bp). The 14 samples were from 7 noncarrier and 6 SANC nostrils, in addition to a microbial community standard that served to validate the library preparation and sequencing resolution. In total, 416 unique species were detected, while about 90 species were common to all noses. A list of the most prevalent genera and species, and the relative abundance and number of colonized nostrils in each group, is provided (Table 1). All seven culture-defined non-SANC nostrils contained *S. aureus* DNA (average relative abundance of 0.05).
abundance, 1.83%), while SANC nostrils averaged 14.53% S. aureus DNA (Fig. 3A and Table 1). Several Staphylococcus and Corynebacterium species appeared to associate with noncarrier noses, including S. capitis, S. epidermidis, S. lugdunensis, Corynebacterium accolens, Corynebacterium segmentosum, and C. pseudodiphtheriticum (Table 1). However, due to the variation between nostrils from different hosts, differences in abundance of these species between persistent and noncarriers did not reach statistical significance. Instead, we evaluated the ratio of various species to S. aureus for individual nostrils in each group. For Acinetobacter spp., whose

TABLE 1 Nasal microbial composition of culture-defined noncarriers and carriers of S. aureusa

| Genus and species          | Noncarrier noses | S. aureus carrier noses |
|---------------------------|------------------|--------------------------|
|                           | Avg. relative abundance (%) | No. of colonized nostrils | Avg. relative abundance (%) | No. of colonized nostrils |
| Staphylococcus            |                  |                          |                          |
| S. aureus                 | 1.829            | 7 of 7                   | 14.528                   | 6 of 6                    |
| S. auricularis            | 0.024            | 3 of 7                   | 0.614                    | 3 of 6                    |
| S. capitis                | 1.936            | 7 of 7                   | 0.439                    | 5 of 6                    |
| S. cohnii                 | 0.064            | 3 of 7                   | 0.006                    | 1 of 6                    |
| S. epidermidis            | 16.669           | 7 of 7                   | 9.140                    | 6 of 6                    |
| S. equorum                | 0.014            | 2 of 7                   | Und                      | 0 of 6                    |
| S. haemolyticus           | 0.050            | 5 of 7                   | 0.461                    | 5 of 6                    |
| S. hominis                | 0.088            | 6 of 7                   | 0.085                    | 6 of 6                    |
| S. lugdunensis            | 1.263            | 5 of 7                   | 0.493                    | 4 of 6                    |
| S. pasteuri               | 0.410            | 7 of 7                   | 0.338                    | 5 of 6                    |
| S. saprophyticus          | 0.006            | 1 of 7                   | 0.009                    | 1 of 6                    |
| S. warneri                | 0.363            | 5 of 7                   | 0.606                    | 5 of 6                    |
| Corynebacterium           |                  |                          |                          |
| C. accolens               | 13.927           | 7 of 7                   | 3.367                    | 6 of 6                    |
| C. segmentosum            | 13.737           | 7 of 7                   | 3.190                    | 6 of 6                    |
| C. aurimucosum            | 6.783            | 7 of 7                   | 12.992                   | 6 of 6                    |
| C. pseudodiphtheriticum   | 1.617            | 7 of 7                   | 0.064                    | 3 of 6                    |
| C. matruchotii            | 1.024            | 7 of 7                   | 0.206                    | 3 of 6                    |
| C. stationis              | 0.134            | 3 of 7                   | 0.006                    | 1 of 6                    |
| C. diphtheriae            | 0.199            | 6 of 7                   | 0.058                    | 6 of 6                    |
| C. kroppenstedtii         | 0.509            | 6 of 7                   | 0.706                    | 5 of 6                    |
| C. simulans               | 0.109            | 6 of 7                   | 0.033                    | 5 of 6                    |
| C. ureiclerivorans        | 0.036            | 5 of 7                   | 0.017                    | 2 of 6                    |
| C. ottidis                | 0.023            | 3 of 7                   | 0.007                    | 1 of 6                    |
| C. fournierii             | 0.020            | 3 of 7                   | 0.006                    | 1 of 6                    |
| C. resistens              | 0.015            | 4 of 7                   | Und                      | 0 of 6                    |
| Cutibacterium acnes       | 9.339            | 7 of 7                   | 8.590                    | 6 of 6                    |
| Cutibacterium granulosum  | 1.113            | 7 of 7                   | 3.233                    | 6 of 6                    |
| Dolosigranulum pigrum     | 2.046            | 7 of 7                   | 0.273                    | 1 of 6                    |
| Citrobacter koseri        | 0.464            | 2 of 7                   | Und                      | 0 of 6                    |
| Klebsiella aerogenes      | 0.174            | 3 of 7                   | 0.006                    | 1 of 6                    |
| Moraxella lincolnii       | 9.296            | 6 of 7                   | 0.071                    | 1 of 6                    |
| Bacillus subtilis         | 0.070            | 3 of 7                   | 1.025                    | 4 of 6                    |
| Streptococcus spp.        | 0.370            | 5 of 7                   | 3.172                    | 6 of 6                    |
| Acinetobacter             |                  |                          |                          |
| A. baumannii              | 0.455            | 6 of 7                   | 1.453                    | 3 of 6                    |
| A. haemolyticus           | 0.008            | 2 of 7                   | 0.013                    | 3 of 6                    |
| A. johnsonii              | 0.097            | 5 of 7                   | 0.208                    | 3 of 6                    |
| A. junii                  | 0.750            | 7 of 7                   | 1.501                    | 5 of 6                    |
| A. lwoffii                | 0.110            | 5 of 7                   | 0.069                    | 5 of 6                    |
| A. pittii                 | 0.148            | 5 of 7                   | 0.074                    | 3 of 6                    |
| A. radioresistens         | 0.006            | 2 of 7                   | 0.020                    | 2 of 6                    |
| A. schindleri             | 0.094            | 5 of 7                   | 0.102                    | 2 of 6                    |
| A. towneri                | 0.013            | 3 of 7                   | 0.037                    | 2 of 6                    |
| A. venetianus             | 0.087            | 3 of 7                   | 0.141                    | 1 of 6                    |
| A. wuhouensis             | 0.114            | 2 of 7                   | 0.028                    | 2 of 6                    |

Und, undetected in all swabs within the indicated group.
relative DNA abundances were low and lacked consensus (only *A. junii* was present in all seven noncarrier nostrils, while no *Acinetobacter* spp. were common to all SANC nostrils [Table 1]), the ratio of total *Acinetobacter* to *S. aureus* was calculated. Six of seven noncarrier noses exhibited an *Acinetobacter*: *S. aureus* ratio of >1, while 5 of 6 carrier ratios were <1 (Fig. 3B; and *P* = 0.026). The carrier nose in which *Acinetobacter* outnumbered *S. aureus* (arrow in Fig. 3B) nearly cleared *S. aureus* 4 days later. The *S. epidermidis*: *S. aureus* and *C. pseudodiphtheriticum* (*C. pseudodiphtheriticum*) to *S. aureus* for the same groups. Error bars represent the mean ± the standard error of the mean (SEM). *P* values are indicated for each comparison (unpaired *t*-tests).

**Survival of *S. aureus* in synthetic nasal medium is potently inhibited by *K. aerogenes, C. koseri, and M. lincolni**i**. We screened 11 nasal bacterial species (17 strains) for inhibitory activity against *S. aureus* using a nutrient background composed of synthetic nasal medium (26) supplemented with 0.1% human serum (here referred to as SNM). Krismer et al. found that *S. aureus* growth in SNM led to gene expression patterns similar to what was observed during *in vivo* colonization (26). All strains (*S. aureus* and putative competitors) were isolated from nostril swabs performed in our laboratory or were nasal isolates purchased from the American Type Culture Collection (ATCC). To reduce growth variations due to differences in isolation procedures or broths used for initial stock preparations and subcultures, all strains were propagated in SNM on a layer of primary nasal epithelium, washed and separated from nasal cells, and stored frozen in single-use aliquots. Five genetically distinct nasal *S. aureus* strains...
from our carrier cohort demonstrated the capacity to proliferate 1,000- to 10,000-fold in SNM in 24 h, while growth rates plateaued at input concentrations greater than $10^5$ CFU/ml (50,000 CFU/0.5 ml in Fig. 4A). The growth rates of *K. aerogenes*, *C. koseri*, *M. lincolnii*, and *S. epidermidis* also exceeded 100-fold using an input of $10^5$ CFU/ml (Fig. 4B), while *Acinetobacter* strains, *C. pseudodiphtheriticum*, and *K. oxytoca* did not thrive in SNM (Fig. 4C).

*K. aerogenes* collected from donor 857, an established noncarrier in our cohort, inhibited the recovery of all 5 *S. aureus* strains by over 99% when incubated as a 50:50 mixture (input = 50,000 CFU of each bacterium; Fig. 4D). To screen which resident nasal species compete with *S. aureus*, mixtures (50,000 *S. aureus* CFU + 50,000 CFU test strain) were incubated, and *S. aureus* recovery was calculated by comparing *S. aureus* CFU on selective agar when *S. aureus* was incubated alone versus with the indicated strain. Individual data points and box and whiskers representing the median, 25th to 75th percentiles, and min/max values are shown for at least 3 experiments using unique nasal strains of *S. aureus*. *P < 0.05; **P < 0.01 versus *S. aureus* incubated alone (100% recovery).**

**FIG 4** Inhibited *S. aureus* survival in nasal medium containing select species from the Gammaproteobacteria class. (A) Multiple unique strains of *S. aureus* were tested for their ability to proliferate in a physiological synthetic nasal medium supplemented with 0.1% human serum (SNM). Error bars represent the mean ± SEM. (B and C) Growth of putative nasal *S. aureus* competitor strains in SNM. Individual data points are shown as open circles. Bars and error bars indicate the mean ± SEM. Colored circles for *A. haemolyticus* indicate the unique strains tested. (D) To screen which resident nasal species compete with *S. aureus*, mixtures (50,000 *S. aureus* CFU + 50,000 CFU test strain) were incubated, and *S. aureus* recovery was calculated by comparing *S. aureus* CFU on selective agar when *S. aureus* was incubated alone versus with the indicated strain.
alone, a mixture containing 10% *K. aerogenes*-D857 and 90% *S. aureus* reduced *S. aureus* recovery to 0.1% (black bars in Fig. 5A), while *K. aerogenes* recovery was 80% for all input ratios tested (red bars in Fig. 5A).

*K. aerogenes*-819-56 inhibited *S. aureus* recovery dose-dependently, as did donor and commercial strains of *C. koseri* and *M. lincolnii* (black bars in Fig. 5B to E). Collectively, these data suggest that *S. aureus* survival in nasal medium is thwarted by relative scant amounts of select Gammaproteobacteria, including *K. aerogenes*, *C. koseri*, and *M. lincolnii*.

**Inhibition of *S. aureus* growth on polarized nasal epithelia by noncarrier-associated Gammaproteobacteria.** To further evaluate promising competitor strains in an environment in which *S. aureus* thrives, bacterial mixtures in SNM were placed on...
primary human nasal epithelial cell (NEC) layers that were polarized on porous Transwell inserts. Without antibiotics in the cells or underlying maintenance medium, which contained growth factors and 5% serum, a $10^4$ CFU/Transwell \textit{S. aureus} inoculum proliferated over 10,000-fold in 24 h (Fig. 6A). When $0.5 \times 10^4$ \textit{S. aureus} CFU/Transwell was incubated with an equivalent amount of test strain, surviving bacteria were determined after 24 h by dilution-plating the apical fluid to selective agars that differentiate \textit{S. aureus} from the \textit{Gammaproteobacteria} of interest. Both \textit{K. aerogenes} strains, \textit{C. koseri}-8832 and \textit{C. koseri}-1791, retained their potent anti-\textit{S. aureus} activity, reducing recovery to <0.1% (Fig. 6B). \textit{C. koseri}-D860 and \textit{M. lincolnii} reduced \textit{S. aureus} recovery to 10% and 2%, respectively. \textit{Acinetobacter haemolyticus} (3 strains), \textit{A. junii}, and \textit{A. schindleri} permitted 30 to 40% \textit{S. aureus} recovery (Fig. 6C). \textit{Acinetobacter johnsonii} and \textit{S. epidermidis} did not limit \textit{S. aureus} recovery (Fig. 6C), although they exhibited over 1,000-fold growth on NEC when incubated without \textit{S. aureus} (not shown). \textit{C. pseudodiphtheriticum} did not grow on NEC and appeared not to impact \textit{S. aureus} survival when coincubated (not shown). The five highly effective strains were retested using an input of 9,000 \textit{S. aureus} CFU/1,000 competitor CFU and inhibited \textit{S. aureus} recovery to <1% (\textit{K. aerogenes}-D857, \textit{C. koseri}-8832), <10% (\textit{K. aerogenes}-816-56, \textit{M. lincolnii}), and <50% (\textit{C. koseri}-D860, Fig. 6D). While \textit{S. aureus} recovery was markedly inhibited, detection of the 5 competitor strains was 70 to 100% compared to when they were cultured on Transwells without \textit{S. aureus} present (Fig. 6E). It should be noted that most \textit{S. aureus} strains damage epithelia when incubated for 24 h without competing bacterial species. After 24 h of

\begin{figure}[h]
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\includegraphics[width=\textwidth]{figure6}
\caption{Select \textit{Gammaproteobacteria} inhibit \textit{S. aureus} survival on nasal cells. \textit{S. aureus} competition assays were performed as described in Fig. 4 and 5 except that incubations occurred in the apical chamber of Transwells containing polarized primary human nasal epithelial cells. (A) Growth of 5 unique strains of \textit{S. aureus} on polarized primary nasal epithelia, presented as input and output (24 h) CFU/well. (B and C) \textit{S. aureus} recovery (%) for incubations of \textit{S. aureus} mixed with an equivalent input of the indicated strain. *, \textit{P} < 0.05; **, \textit{P} < 0.01 versus \textit{S. aureus} alone (100% recovery). (D) \textit{S. aureus} recovery (%) for incubations containing 90% \textit{S. aureus} + 10% indicated strain. ***, \textit{P} < 0.001; ****, \textit{P} < 0.0001 versus \textit{S. aureus} incubated alone (100% recovery). (E) Percent recovery of competitor strains from the 90% \textit{S. aureus}/10% test strain mixtures presented in panel D. For all panels, bars and error bars represent the mean ± SEM.}
\end{figure}
incubation in the presented experiments, visual inspection of the Transwells prior to collection of bacteria (which disrupted the cell layers) showed that cells incubated with S. aureus alone were heavily damaged. S. aureus recovery from the basolateral chamber occurred only in wells in which S. aureus was inoculated alone. These experiments suggested that K. aerogenes, C. koseri, and M. lincolnii limit S. aureus survival in the nasal mucosa.

**K. aerogenes, C. koseri, and M. lincolnii inhibit S. aureus survival in nasal secretions.** To test for activity of K. aerogenes, C. koseri, and M. lincolnii against nasal S. aureus strains in an environment more physiological than SNM, assays were performed similarly to above except that a medium background of human nasal fluid (NF) was used. Whole nasal fluid (no dilution during collection) was processed briefly to eliminate endogenous bacterial CFU, and an NF pool was prepared by mixing equal volumes of processed NF from 10 healthy donors. Treatment wells (0.1 ml NF) were incubated alone (to confirm the absence of CFU after processing), with $10^4$ S. aureus CFU, or with 5,000 S. aureus CFU mixed with 5,000 CFU of test strain. For three different strains of S. aureus (2 nasal isolates and USA300), growth in NF exceeded 1,000-fold in 24 h (Fig. 7). K. aerogenes-D857 and C. koseri-8832 reduced S. aureus recovery to <1%, and K. aerogenes-816-56, M. lincolnii, and C. koseri-D860 reduced S. aureus recovery to <10% (Fig. 7A). Notably, these strains showed the capacity to survive and proliferate in NF while attenuating S. aureus (Fig. 7B). Collectively, the presented experiments have identified resident nasal species that limit S. aureus survival and detection by routine nasal sampling methods. K. aerogenes, C. koseri, and M. lincolnii were next tested for secreted factors that inhibit S. aureus growth.

**Conditioned medium from NEC plus K. aerogenes and NEC plus M. lincolnii cocultures exhibit anti-S. aureus activity mediated by heat-stable <30-kDa proteins.** Conditioned medium (CM) was collected from confluent primary NECs incubated alone or with individual bacterial strains for 24 h in SNM. Prior to experiments with S. aureus, CMs were clarified by centrifugation, followed by 0.2 μm filtration, confirmed free of bacterial CFU by agar plating, and frozen in single-use aliquots. The growth of three different strains of S. aureus (nasal 547-14 sequence type 5 [ST5], nasal 512 [ST30], USA300 [ST8]) was evaluated in freshly thawed CM or fresh SNM ($10^4$ CFU/0.1 ml) after incubation for 24 h. Initial S. aureus inocula (e.g., CFU/well at time 0 h) were confirmed by agar plating and subsequent CFU enumeration. S. aureus survival at 24 h was determined by plating multiple dilutions of each fluid and calculating the output CFU per treatment well (Fig. 8). CM from NEC coincubations with K. aerogenes-D857, K. aerogenes-816-56, and M. lincolnii inhibited S. aureus detection by 2 to 3 logs compared to

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**FIG 7** K. aerogenes, C. koseri, and M. lincolnii inhibit S. aureus survival in nasal secretions. Three unique strains of S. aureus (2 nasal isolates and USA300) were incubated alone or with an equivalent input of the indicated strain in undiluted human nasal fluid which was processed briefly to prevent growth of endogenous microbes. (A) S. aureus recovery (%) for the presented incubations. Bars represent the mean ± SEM; n=3; ***, P<0.0001 versus S. aureus alone (100% recovery). (B) Fold growth in nasal fluid for S. aureus alone (3 strains, black circles) or the indicated competitor (open circles) when mixed with S. aureus. Error bars represent the mean ± SEM.
**S. aureus** recovered from plain SNM, CM from untreated NEC, or CM from NEC plus **S. epidermidis** coincubation (Fig. 8A). CM from **C. koseri**- and **S. aureus**-NEC CMs inhibited **S. aureus** recovery by about 10-fold. This trend was conserved when CMs were heat-treated (95°C/5 min) prior to incubation with **S. aureus** (Fig. 8B), suggesting the presence of heat-stable anti-**S. aureus** proteins or peptides. Proteinase K treatment inhibited growth of **S. aureus** by heat-stable <30-kDa proteins found in CM from **K. aerogenes**- and **M. lincolnii**-treated NEC. **S. aureus** (~10⁴ CFU) was incubated 24 h in synthetic nasal medium (SNM) or conditioned medium (CM) from primary nasal epithelial cells (NEC) incubated alone or with the indicated bacteria in SNM, SE, **S. epidermidis**; Ka, **K. aerogenes**; CK, **C. koseri**; M, **M. lincolnii**; SA, **S. aureus**. Collected CM was centrifuged and filter-sterilized and confirmed to be devoid of bacterial CFU prior to experimentation. (A to F) Growth of **S. aureus** is displayed as CFU determined by agar plating for raw CM (A), 95°C/5 min-treated CM (B), proteinase K-treated CM (C), vehicle (storage solution for proteinase K)-treated CM (D), <30-kDa protein fraction of CM (E), and >30-kDa protein fraction of CM (F). Error bars indicate the mean ± SEM for each treatment group. Each circle represents a unique strain of **S. aureus**. *, P < 0.05 versus **S. aureus** recovery from SNM.
abrogated the effect of *K. aerogenes-* and *M. lincolnii*-NEC CMs (Fig. 8C), while vehicle treatment of CMs with proteinase K storage solution preserved the anti-*S. aureus* activity (Fig. 8D). Last, we tested whether <30 kDa proteins mediate the observed anti-*S. aureus* activity by processing CMs through 30 kDa molecular weight cutoff filters. Figure 8E shows that the anti-*S. aureus* potency of *K. aerogenes-* and *M. lincolnii*-NEC CMs is retained in the <30-kDa fraction. *S. aureus* growth and viability in the >30-kDa CM fractions were similar for all bacterium-NEC combinations (Fig. 8F). CM from *Klebsiella* and *Moraxella* cultures, incubated in SNM but without NEC, markedly inhibited *S. aureus recovery compared to uninoculated SNM or CM from NEC cultures (not shown). These data suggest that secreted <30-kDa proteins from nasal *K. aerogenes* and *M. lincolnii* limit the detection of nasal *S. aureus*.

**DISCUSSION**

The nasal cavity filters thousands of liters of air per day, and the nasal mucosa secretes about a liter of fluid per day (37). Passage of inhaled particles or microorganisms up and out of the nasal cavity or lungs is achieved when mucous-trapped inhaledants are moved by epithelial cilia toward the throat, where they are swallowed or expelled by coughing. The anterior nares (nostrils) represent a transition zone from skin to the airway mucosa and are not part of this mucociliary escalator. Thus, the resident microflora of this anatomical region, combined with microbicidal substances, help shape the nasal mucosal immune ecosystem of each host. SANC presents an obvious risk factor for autologous infection, as *S. aureus* is one of the most common pathogens isolated from skin and soft tissue wounds and surgical sites, and nasally carried strains are usually genetically indistinguishable from strains isolated from infection sites (7, 38, 39). Intermittent and persistent carriers acquire and lose *S. aureus* strains at similar frequencies (5, 40), but persistent carriers possess a higher nasal *S. aureus* load than intermittent carriers. This brings into question whether intermittent and noncarriers truly exist or just carry *S. aureus* below culture detection limits (23, 41). To date, it is not understood why some healthy humans carry detectable levels of nasal *S. aureus* persistently, intermittently, or seemingly not at all. Metabolomic analysis of nasal secretions showed low levels of potential nutrients and no appreciable differences between SANC and noncarrier secretions (26), and few studies have explored specifically the nasal microbiomes of established carriers versus noncarriers. The positive correlation between nasal *S. aureus* load and ex vivo growth of *S. aureus* strains in matched nasal fluids from SANC subjects (Fig. 1) suggested that *S. aureus* CFU detection might relate to the composition and quantity of microbiome-derived soluble proteins contained in nasal fluids. We therefore set out to determine species-level differences in the microbial composition of established SANC and non-SANC nostrils and to test noncarrier-associated microbial community members for activity against *S. aureus* in an environment mimicking the nasal milieu.

Our multipronged approach included 16S amplicon sequencing of 65 swab samples collected from established noncarrier, intermittent, and persistent SANC nostrils; shotgun metagenomic sequencing of DNA collected freshly from 7 established noncarrier swabs and 6 persistent SANC swabs; and culture-based isolation and identification of nasal species from non-SANC noses. Shotgun DNA sequencing afforded species- and strain-level resolution, and all presumed non-SANC subjects (based on culture results) had detectable *S. aureus* DNA, while one noncarrier exhibited 11% *S. aureus* DNA (relative abundance; Fig. 3A). Culture *S. aureus*-positive noses carried *S. aureus* ranging in relative abundance between 0.7% and 33%, demonstrating that some people do not harbor high levels of *S. aureus* DNA but still swab positive persistently, while some presumed “noncarriers” host abundant *S. aureus* DNA without exhibiting CFU. Others have reported nasal *S. aureus* DNA to be nearly always present (42, 43), and all people are thought to possess anti-*S. aureus* antibodies (44). Taken together, these findings support the concept that people are consistently exposed to *S. aureus* in everyday life and that SANC likely represents a microbial shift condition that can be modulated with
probiotics or improved topical anti-\textit{S. aureus} biologicals once the key influential species are identified.

The \textit{Gammaproteobacteria} genera \textit{Citrobacter} and \textit{Moraxella} were detected using 16S amplicon sequencing and associated with some non-SANC nostrils (Fig. 2A), and both \textit{C. koseri} and \textit{M. lincolnii} were found in greater relative abundance in noncarrier nostrils when assessed by shotgun sequencing (Table 1). \textit{K. aerogenes} was not as abundant in nostril swabs as \textit{M. lincolnii} or \textit{C. koseri}, but it was associated more with noncarrier noses (Table 1). Notably, we were able to isolate \textit{C. koseri} and \textit{K. aerogenes} from noncarrier but not carrier noses. These strains as well as purchased nasal strains of \textit{C. koseri}, \textit{K. aerogenes}, and \textit{M. lincolnii} proliferated upward of 1,000-fold in SNM, indicating that they can adapt to the nutrient-limited environment of the anterior nares (Fig. 4). Collectively, the finding that these species strongly inhibited \textit{S. aureus} recovery even when the \textit{S. aureus:competitor} input ratio was 90:10 and the test environment (nasal epithelia or nasal fluid) supported rampant \textit{S. aureus} growth when no competitor was added (Fig. 5 and 7) suggests that even relative scant amounts of these species might prevent detection of nasal \textit{S. aureus} by standard swabbing and culture-based procedures. Likewise, the average competitor: \textit{S. aureus} DNA ratios in established noncarriers were 1:10.5 for \textit{K. aerogenes}, 1:4 for \textit{C. koseri}, and 5:1 for \textit{M. lincolnii} (Table 1), thus providing a plausible explanation for why these “noncarrier” subjects were swabbed weekly for months to years without detection of a single \textit{S. aureus} colony. A study of the nasal microbiomes of chronic rhinitis patients receiving intranasal steroids demonstrated that suppression of \textit{Moraxella} spp. was associated with increased abundance of \textit{Staphylococcus} spp. (45). Vaginal microbial profiling showed an inverse relationship between \textit{Klebsiella} spp. prevalence and \textit{S. aureus} culture positivity (46). Correspondingly, Liu et al. defined seven major nasal community state types (CSTs) and discovered that nasal microbiomes classified as CST2 [\textit{Enterobacteriaceae} e.g., \textit{Klebsiella} and \textit{Citrobacter} prevalent] and CST6 (\textit{Moraxella} spp. prevalent) were not likely to contain \textit{S. aureus} (28). Since CM derived from \textit{K. aerogenes} and \textit{M. lincolnii} cultures inhibited \textit{S. aureus} while CM from \textit{C. koseri} did not (Fig. 8), it is apparent that this class of \textit{Gammaproteobacteria} possesses multiple unique mechanisms for limiting survival of nasal \textit{S. aureus} that warrant further investigation.

Implementation of \textit{K. aerogenes} or \textit{C. koseri} as nasal therapeutics for \textit{S. aureus} decolonization might be challenging, as these species are known opportunistic pathogens. While not as dreaded as \textit{K. pneumonia}, \textit{K. aerogenes} is associated with urinary tract and bloodstream infections in vulnerable populations and is capable of acquiring antibiotic resistance (47). \textit{C. koseri} is common to the intestinal tract, and though seldom lethal, is known to cause a variety of infections in humans. On the other hand, we isolated \textit{K. aerogenes}, \textit{K. oxytoca}, and \textit{C. koseri} from nasal swabs collected from healthy volunteers. We noted that topical administration of \textit{K. aerogenes}, \textit{M. lincolnii}, and \textit{C. koseri} to polarized primary nasal epithelia resulted in less barrier disruption than administration of \textit{S. aureus}, but our assays were designed for liberation and enumeration of CFU instead of host cell signaling and toxicity studies. A comprehensive analysis of the effects of these species on nasal tissue (e.g., inflammatory markers, epithelial barrier integrity, chemotactic activity) is needed. In the meantime, identification and characterization of anti-\textit{S. aureus} protein(s) from these species should include plans for chemical (bacterium-free) synthesis of active peptides as well as safety, stability, and functional testing in nasal tissue and against a variety of resident nasal bacteria. \textit{M. lincolnii} has not been well characterized to date, but it is considered a normal resident of the upper respiratory tract. High abundance of nasal \textit{Moraxella} spp. is associated with healthy young children and low rates of respiratory infections (48), and a longstanding healthy \textit{S. aureus} noncarrier in our cohort was classified as nasal \textit{M. lincolnii} prevalent (>60% relative bacterial DNA abundance). Thus, \textit{M. lincolnii} represents a promising candidate for preclinical testing as a natural \textit{S. aureus} competitor on mucosal surfaces and perhaps skin lesions. Future studies should also test whether \textit{S. aureus} can develop resistance to the \textit{Gammaproteobacteria}-derived peptide(s) and whether there is a
fitness and/or virulence cost. Given that few novel anti-\textit{S. aureus} antibiotics have been brought to market over the past several decades, while \textit{S. aureus} infections continue to burden health care systems worldwide, a renewed effort to exploit commensal species and their peptide products may prove worthwhile.

Numerous reports have indicated roles for \textit{Corynebacterium} spp. and \textit{S. epidermidis} in favoring the non-SANC state, and mechanisms of interaction between these commensals and \textit{S. aureus} have been described (30, 33, 49–51). We found that few obvious distinctions between non-SANC, intermittent, and persistent SANC nostrils were indicated by 16S amplicon sequencing aside from the expected trend that \textit{Staphylococcus} abundance appeared to be inversely proportional to \textit{Corynebacterium} abundance in some individuals who were sampled multiple times (underlined stacked bars in Fig. S1). Interestingly, however, wide ranges of \textit{Staphylococcus} and \textit{Corynebacterium} abundance were revealed in both non-SANC and persistent SANC individuals; a persistent SANC nostril contained \(<10\% \textit{Staphylococcus} and \textit{Corynebacterium} relative abundance, while a noncarrier nostril contained \(>80\% \textit{Staphylococcus} relative abundance (Fig. 2A and Fig. S1). These observations suggest that the competitive relationship between \textit{Corynebacterium} spp. and nasal \textit{S. aureus} is highly complex and not yet well understood.

Shotgun metagenomic sequencing revealed that \textit{Corynebacterium} spp. (particularly \textit{C. accolens}, \textit{C. segmentosum}, and \textit{C. aurimucosum}) and \textit{S. epidermidis} were abundant in all noses (Table 1). A potential limitation of our study is that, due to cost considerations, we performed shotgun sequencing on only 13 swab samples (e.g., single HiSeq lane). However, the species-level data uncovered several potential distinctions between persistent- and non-SANC that are in line with recent reports from other groups or shed light on unanswered questions. Noncarrier noses were associated with an abundance of \textit{C. accolens} and \textit{C. segmentosum} compared to \(>10\) other \textit{Corynebacterium} spp., while SANC nostrils were colonized mainly by \textit{C. aurimucosum} (Table 1). Nasal \textit{C. accolens} was determined to be more prevalent in the absence of \textit{S. aureus}, although during longitudinal sampling, it was unclear if its presence impacted future acquisition of nasal \textit{S. aureus} (52). \textit{C. aurimucosum} showed mixed results against 3 strains of \textit{S. aureus} in agar-based zone of clearance assays, suggesting mainly contact-independent bacteriostatic interaction (31). \textit{C. pseudodiphtheriticum} interaction with \textit{S. aureus} may be bactericidal or diminish quorum sensing and fitness (33, 49). Our data revealed that \textit{C. pseudodiphtheriticum} was present in all noncarrier nostrils but only half of carrier noses, supporting the view that this species reduces the rate of detectable SANC. On the other hand, \textit{C. pseudodiphtheriticum} did not thrive in SNM or inhibit \textit{S. aureus} in the nutrient background of SNM (Fig. 4) or SNM placed on the apical surface of polarized NEC. This is consistent with the report that \textit{C. pseudodiphtheriticum}’s niche is the middle meatus/sphenoid recess and not the anterior nares (34). Nasal \textit{S. epidermidis} grew in SNM but failed to limit \textit{S. aureus} growth (Fig. 4 and 6), supporting Liu et al. (28) and our previous work (15) finding that \textit{S. epidermidis} coexists with \textit{S. aureus} in the nose. Indeed, we observed CST3 (\textit{S. epidermidis} prevalent) and CST5 (\textit{Corynebacterium} spp. prevalent) in both the non-SANC and persistent SANC groups, while CST6 (\textit{Moraxella} spp. prevalent) was observed only in a long-term “never” \textit{S. aureus}-positive subject. Our shotgun sequencing data also corroborated the findings that \textit{S. lugdunensis} and \textit{Dolosigranulum pigrum} levels trend higher in non-SANC nostrils (28, 32). \textit{S. lugdunensis} produces a peptide antibiotic, “lugdunin,” that inhibited SANC in cotton rats, and a survey of hospitalized patients found that although nasal carriage of \textit{S. lugdunensis} was \(>3\)-fold less common than SANC, only 1 of 17 \textit{S. lugdunensis}-positive noses was \textit{S. aureus}-positive (32). Future studies aimed at defining the anti-\textit{S. aureus} properties of \textit{D. pigrum} should be undertaken.

It was surprising that only the ratio of \textit{Acinetobacter} spp. to \textit{S. aureus} was significantly different between the \textit{S. aureus} culture-negative versus \textit{S. aureus} culture-positive swabs (Fig. 3B), considering that overall abundance of \textit{Acinetobacter} was considerably
lower than that of Corynebacterium, non-aureus Staphylococcus, Cutibacterium, and Dolosigranulum (Table 1). Six of seven noncarrier noses exhibited an Acinetobacter:S. aureus ratio of >1, while five of six carrier ratios were <1. Interestingly, the carrier nose in which Acinetobacter outnumbered S. aureus (arrow in Fig. 3B) nearly cleared S. aureus according to a swab sample collected 4 days later. To date, only Acinetobacter baumannii has received considerable attention in airway studies due to its frequent association with pneumonia and its propensity for gaining antibiotic resistance. We failed to cultivate Acinetobacter strains from fresh nostril swabs and, thus, purchased them from ATCC and prepared subcultures and then sequenced the 16S gene to reconfirm their identities. We noted that A. baumannii genetic sequences provide the basis for sequence mapping of less well-characterized Acinetobacter spp. and were always included in BLAST results. A recent screen of wound and nasal isolates determined that select strains identified by 16S sequencing as A. baumannii possess contact-dependent activity against S. aureus (31). Our data also suggest contact-dependent S. aureus inhibition, as A. haemolyticus, A. junii, and A. schindleri all showed some anti-S. aureus activity when incubated with diverse S. aureus strains on polarized nasal epithelia (Fig. 6C), while CM collected from these species did not prevent S. aureus growth (data not shown). We conclude that Acinetobacter spp. warrant further investigation as S. aureus effectors, as they may sequester a vital S. aureus nutrient or achieve bactericidal activity via direct contact. A recent report of a <15% SANC rate in a long-term care home in which residents were heavily colonized with Acinetobacter (S3) provides support for this concept.

In conclusion, we discovered members of the Gammaproteobacteria class that reside in S. aureus-negative noses and possess anti-S. aureus properties. K. aerogenes, C. koseri, and M. lincolnii inhibited S. aureus growth and recovery by over 90% in physiologically relevant assays containing synthetic nasal medium, polarized primary nasal epithelia, and nasal secretions. Select Acinetobacter species, including haemolyticus, junii, and schindleri, were found to be slow growing on nasal epithelia compared to S. aureus but still inhibited S. aureus recovery by over 50%. By comparing the conditioned media from bacterium plus epithelium cocultures, we further revealed that K. aerogenes and M. lincolnii secrete <30-kDa protein(s) that inhibit S. aureus, while C. koseri and Acinetobacter spp. activity against S. aureus appeared to be contact-dependent. Taken together, the presented data provide an explanation for why some people appear to be noncarriers of S. aureus even though all noses contain S. aureus DNA. Further investigation is needed to define the mechanisms of anti-S. aureus activity, but this study introduces the concept that acquisition and loss of detectable SANC might result from fluctuations in the nasal microbiota. Better understanding of nasal microbial community dynamics and how the nasal microbiome controls S. aureus viability and growth is greatly needed in order to design effective decolonization strategies.

MATERIALS AND METHODS

Ethics statement. A protocol for recruitment of healthy participants and collection of nasal swabs and nasal fluid was approved by the Institutional Review Board of the University of Central Florida. Participants reported no adverse effects.

Sample collection. The Staphylococcus aureus strains used in this study were collected from healthy, nasally colonized subjects by our laboratory and genotyped by multilocus sequence typing (MLST) and spa typing as described previously (5, 54). Specifically, the strains were 547-14 (ST5, t688), 512 (ST30, t012), 528-11 (ST8, t008), 594-5 (ST188, t037), and 713-4 (ST5, t548). USA300 (ST8, t008, multidrug resistant) was acquired from NARSA (now www.beiresources.org). These represented a cross section of the most common sequence types described in our large longitudinal study of over 100 nasal S. aureus carriers (5). In an effort to isolate and identify prevalent nasal strains from established non-S. aureus carriers (>3 months of weekly S. aureus-negative swabs) or intermittent carriers who swabbed S. aureus-negative 1 to 3 days prior, participants sampled each nostril by rotating a sterile polyester-tipped swab around the anterior vestibule 10 times. Swab tips were vortexed in 0.5 ml TSBS (Bacto tryptic soy broth; Becton, Dickinson [BD], Franklin Lakes, NJ) to liberate the microbes. From each 0.5-ml mixture of microbes and broth, 0.1 ml was spread onto each of the following agars: BD CHROMagar S. aureus (Fisher Scientific; catalog no. 14-432-41) for identification of S. aureus, nonselective TSA II/5% sheep blood agar (Fisher Scientific; catalog no. B21261X), MacConkey II (Fisher Scientific; catalog no. B21270X) for selection of mainly Gram-negative species, Leeds agar (Hardy Diagnostics; catalog no. G261), or Hoyle’s tellurite agar...
Nasal Microbes Inhibit *S. aureus*  

(1 liter; 10 g Lab Lemco beef extract, 10 g peptone, 5 g NaCl, 15 g agar, 7% horse blood, 0.35 g potassium tellurite; all components from Fisher). On a few collection days early in the study, plating to nutrient agar, plate count agar, and plain TSA occurred instead of plating to MacConkey II, Leeds, and Hoyle's agars. These nonselective agars yielded mostly *Staphylococcus* CFU. Participants’ nasal fluids were collected by suction catheter as described previously (15, 21), and stored at −80°C immediately.

**Nasal bacterial strain details and stock preparation.** Colonies plucked from agar plates were cultured in 4 ml TSB overnight (37°C/250 rpm). The next day, 1 ml was supplemented with glycerol to 15% and stored at −80°C, and the rest was centrifuged to collect bacteria prior to DNA extraction (Qiagen catalog no. 12224-250; UltraClean microbial DNA isolation kit). 16S PCR was performed (standard 27F-1493R primers), verified by gel electrophoresis, and then sent for Sanger sequencing (27F, 341F, 1493R primers) at Eton Biosciences (Research Triangle Park, NC). Sequencing files were analyzed using Mega7.0.20 software to align and concatenate the PCR product sequences into the full-length 16S gene.

Illumina adapter sequences (Nextera XT index primers) at Eton BioSciences (Research Triangle Park, NC). Sequencing was performed on TapeStation 4200 (Agilent) and sent to the University of Florida Biotechnology Research (ICBR) for MiSeq sequencing (2

**Nasal swab samples were tested for functional antibiotic resistance by performing turbidity (growth) assays (55). Isolates were diluted to 10⁶ CFU/90 μl in Mueller-Hinton broth (Millipore Sigma, St. Louis, MO) containing 5% sucrose and loaded onto a 96-well (96W) culture plate. Then, 10 μl of 2-fold serial dilutions of either mupirocin, tobramycin, gentamicin, penicillin-streptomycin, or vehicle (volume-matched dimethyl sulfoxide (DMSO) or phosphate-buffered saline (PBS)) were added to the diluted *S. aureus* so that the final concentration of antibiotic ranged from 1 to 100 μg/ml. Each culture plate was covered with ThermoSeal A film (Excel Scientific, Inc., catalog no. TSA-100) and placed into a SpectraMax 190 microplate reader (Molecular Devices, Sunnyvale, CA) programmed for a 16-h kinetic assay at 37°C. Turbidity measurements (optical density [OD] at 550 nm) were taken every 5 min following 15 sec of agitation. Growth curves (OD at 550 nm plotted against time) were generated for all wells, and both the input and 16-h incubations were plated on TSA II/5% sheep blood agar for enumeration. All strains grew to 10⁷ to 10⁸ CFU/ml in antibiotic-free medium. All *S. aureus* strains were killed in the presence of 1 μg/ml mupirocin, while all *Proteobacteria* tested were mupirocin resistant. All nasal strains tested were killed in the presence of 1 to 10 μg/ml tobramycin, ≥10 μg/ml gentamicin, and penicillin (100 IU/ml)-streptomycin (100 μg/ml).

**Next-generation sequencing (NGS) of nasal swabs.** Nasal swab samples were first assayed for taxonomic composition by sequencing the V3-V4 variable region of the 16S rRNA gene (primers 341F and 785R [56]) on Illumina’s MiSeq platform. In brief, DNA was extracted from nasal swabs using the ZymoBIOMICS DNA microprep kit (Zymo Research Corp., Irvine, CA). PCR was carried out according to Illumina’s instructions (16S metagenomic sequencing library preparation protocol) using an amplicon primer with specified overhangs, and bead cleanup was followed by indexing with uniquely barcoded Illumina adapter sequences (Nextera XT index primers). After a second bead purification of indexed amplicons, samples were quantified (Quant-IT PicoGreen; Molecular Probes, Inc., Eugene, OR) and pooled according to equimolar content. The final sequencing pool was checked for quality using TapeStation 4200 (Agilent) and sent to the University of Florida’s Interdisciplinary Center for Biotechnology Research (ICBR) for MiSeq sequencing (2 × 300 bp). The sequence data were processed using the USEARCH/UPARSE program suite (57, 58) to identify operational taxonomic units (OTUs), and OTU taxonomic assignments were made using a naive Bayesian classifier implemented in mothur (59, 60). OTUs with the same genus assignment were combined to identify the relative abundances of bacterial genera in the nasal microbiomes of a cohort of healthy human volunteers determined to be either long-term (“persistent”), intermittent, or noncarriers of nasal *S. aureus* (n = 65). The OTU data were used to calculate alpha diversity (Shannon diversity index). The genus-level data were used to cluster the samples based on the DMM framework (35). Since species- and strain-level differences cannot be inferred by studying 16S rRNA genes alone, we performed shotgun metagenomics analysis of swab samples from carriers and noncarriers to garner species/strain-
level details. Fresh nasal swabs were collected from volunteers whose SANC status had been established for at least 3 months and was confirmed by culture 1 to 3 days prior, and microbial DNA was enriched by depleting cell-free and mammalian DNA prior to microbial DNA extraction (MoYsis Basic5 microbial DNA enrichment kit; Oasis Diagnostics Corp.). Nextera XT libraries (Illumina, Inc., San Diego, CA) were prepared and indexed according to company instructions, sizes and quality verified using an Agilent 4200 TapeStation system and high sensitivity (HS) D1000 reagents, and pooled. Altogether, 14 samples (7 noncarriers/6 carriers/1 microbial community standard [Zymo Research Corp., Irvine, CA]) were sequenced (HiSeq 2 × 150-bp/dual index/1 lane; Genewiz, Inc.), yielding 462 million reads with a mean quality score of 36.72. After trimming of adapter sequences and low-quality bases, the reads were mapped to the human genome. Unmapped reads from this step were subsequently mapped against a collection of >10,000 microbial genomes (NCBI RefSeq).

This read mapping information was used to estimate relative abundances of genomes (strains) in each sample using an expectation maximization algorithm (61, 62). The most abundant genomes (>0.01 genome abundance) accounted for >150 species, ~90 of which were common to most nasal swabs. For comparisons between noses in which certain species were not detected, the relative abundance was set at 0.005.

Preparation of simulated nasal medium and first-pass *S. aureus* competitor screening. Synthetic nasal medium (SNM) was prepared as described for “SNM3” by Krismer et al. (26). All amino acids, organic acids, trace elements, cofactors, and 2,2'-bipyridine were purchased from Millipore Sigma (St. Louis, MO). Sodium phosphate, potassium phosphate, sodium chloride, potassium chloride, and magnesium sulfate were purchased from Fisher Scientific (part of Thermo Fisher, Waltham, MA). Stock solutions of the components were prepared at 40 to 2,000-fold and stored at 4°C in the dark for ease of use and standardization of individual batches. SNM was demonstrated to mimic the environment of the nasal mucosa and support growth and gene expression observed in human nasal *S. aureus* colonization (26). For all presented experiments, we utilized SNM supplemented with 0.1% (vol/vol) human serum (GeminolBio, West Sacramento, CA). In testing the growth capacities of various nasal microbial community inhabitants in SNM/0.1% serum, inputs ranging from 10,000 to 100,000 CFU per 0.5 ml were incubated 24 h in 24W tissue culture plates at 37°C/5% CO2.

All *Staphylococcus* strains and most *Klebsiella*, *Citrobacter*, and *Moraxella* strains proliferated >100-fold at these initial densities, and the first round of competition assays tested 50,000 CFU *S. aureus* versus 50,000 CFU competitor in 0.5 ml SNM/0.1% serum. When different ratios were used, the total bacterial input was held constant at 100,000 CFU; 90:10 was defined as 90,000 CFU *S. aureus* versus 10,000 CFU competitor, 70:30 meant 70,000 CFU *S. aureus* versus 30,000 CFU competitor, etc. In every experiment, a portion of the prepared bacterial solutions was plated on *S. aureus*, *Klebsiella*, and TSA II/5% blood to confirm the input concentration and colony phenotypes. At 24 h, treatment wells were mixed by pipetting up and down and were plated in serial dilutions to the same agars as follows: CHROMagar *S. aureus* to enumerate *S. aureus* CFU, MacConkey II to enumerate *Klebsiella*, *Citrobacter*, *Moraxella*, and select *Acinetobacter* CFU, and TSA II/5% blood agar for visualization of all strains’ CFU.

Air-liquid interface primary nasal epithelial cell culture. Human primary nasal epithelial cells (NEC; catalog no. T4014) and Prigrow I medium (catalog no. TM001) were purchased from Applied Biological Materials, Inc. (Richmond, BC, Canada). Cells were maintained and expanded on collagen-coated tissue culture plates using a medium of Prigrow I supplemented with 5% fetal bovine serum (GeminiBio, West Sacramento, CA). In testing the growth capacities of various nasal microbial community inhabitants in SNM/0.1% serum, inputs ranging from 10,000 to 100,000 CFU per 0.5 ml were incubated 24 h in 24W tissue culture dishes at 37°C/5% CO2. All *Staphylococcus* strains and most *Klebsiella*, *Citrobacter*, and *Moraxella* strains proliferated >100-fold at these initial densities, and the first round of competition assays tested 50,000 CFU *S. aureus* versus 50,000 CFU competitor in 0.5 ml SNM/0.1% serum. When different ratios were used, the total bacterial input was held constant at 100,000 CFU; 90:10 was defined as 90,000 CFU *S. aureus* versus 10,000 CFU competitor, 70:30 meant 70,000 CFU *S. aureus* versus 30,000 CFU competitor, etc. In every experiment, a portion of the prepared bacterial solutions was plated on *S. aureus*, *Klebsiella*, and TSA II/5% blood to confirm the input concentration and colony phenotypes. At 24 h, treatment wells were mixed by pipetting up and down and were plated in serial dilutions to the same agars as follows: CHROMagar *S. aureus* to enumerate *S. aureus* CFU, MacConkey II to enumerate *Klebsiella*, *Citrobacter*, *Moraxella*, and select *Acinetobacter* CFU, and TSA II/5% blood agar for visualization of all strains’ CFU.
floating cells) were transferred to microtubes, and the remaining cells were vigorously washed and transferred to the same collection tube using ice-cold Hanks’ balanced salt solution (HBSS). Underlays were collected in their own microtubes and stored on ice until agar plating for bacterial enumeration. Apical fluid-rinse mixtures were centrifuged at 300 × g (4°C, 5 min) to separate epithelial cellular debris from bacteria. Bacteria were then collected by centrifugation at 10,000 × g (4°C, 5 min), suspended in HBSS, and dilution-plated to various agar formulations—CHROMagar S. aureus, TSA II/5% sheep blood, and MacConkey II.

**Nasal fluid processing and incubation with S. aureus and competitor strains.** Since collected nasal fluids (NF) were immediately frozen as described above, processing occurred at the time of first thaw. Fluids were pulse-sonicated (1 sec × 30) on wet ice using power level 2 (Fisher Scientific sonic dismembrator model 100), clarified by centrifugation (13,000 rpm/5 min/4°C), and then incubated at 55°C for 20 min to prevent the growth of endogenous bacteria. This method was determined to be preferable to filtration, as 0.2 μm filtration of NF was determined to result in 20% loss of total protein. Compared to heat treatment at 95°C for 5 min, the 55°C/20 min method was also found to preserve protein composition, as evidenced from gel analysis. NF was confirmed free of live bacteria by direct plating to TSA II/5% blood agar and observing no visible CFU after 48 h (37°C bacterial oven). To test for activity of *C. koseri*, *K. aerogenes*, and *M. lincolnii* against nasal *S. aureus* strains in NF, processed NF was pooled from 10 donors in order to create a volume that would enable sufficient experimental replicates. Each bacterial strain was diluted in pooled NF to achieve an incubation volume of 100 μl containing 5,000 CFU of each bacterium pair (e.g., *C. koseri* versus *S. aureus*) or 10,000 CFU of *S. aureus* alone. Mixtures were inoculated at 37°C/5% CO2 in 96W tissue culture plates for 24 h before dilution plating to CHROMagar *S. aureus* or MacConkey II to enumerate *S. aureus* versus *Citrobacter*, *Klebsiella*, or *Moraxella* colonies.

**Preparation and processing of conditioned medium for evaluation of anti-*S. aureus* effectors.** To prepare conditioned medium (CM) from bacteria plus NEC cocultures, 100-mm tissue culture dishes containing a confluent monolayer of primary NEC were treated with 8 ml/dish of SNM/0.1% human serum containing ~4 × 106 CFU (multiplicity of infection [MOI] ~1) of the bacterium of interest (e.g., *C. koseri*, *K. aerogenes*, *M. lincolnii*, etc.). Immediately after the treated dishes were placed at 37°C/5% CO2, serial dilutions of bacteria were plated on agar to confirm the inoculum. After 24 h, each dish was scraped with a cell lifter, and the combined mixture of cells, bacteria, and supernatant was transferred to conical tubes. After centrifugation at 300 × g (4°C, 5 min), which pelleted NEC, the resulting supernatant was centrifuged again at 10,000 × g (4°C, 5 min) to collect bacteria. This clarified supernatant was then passed through a 0.2-μm polyethersulfone (PES) syringe filter (Coming; catalog no. 431229) to remove residual bacteria, and the resulting CM was tested for bacterial growth by direct plating to TSA II/0.5% blood agar plates and stored in aliquots at −80°C until use. CM containing bacteria alone was made in exactly the same way, using 100-mm tissue culture dishes but no epithelial cells. Uninoculated NEC (no bacteria) CM was made in parallel with NEC plus bacteria coinoculations. Anti-*S. aureus* activity of individual CM preparations was evaluated by adding 10 μl (10,000 CFU) of freshly thawed and diluted *S. aureus* to 90 μl of freshly thawed CM in a 96W tissue culture dish and incubating it at 37°C/5% CO2 for 24 h. Bacteria were enumerated by plating serial dilutions on agar plates, and fold growth was calculated (Tfinal/Tinoculum).

To characterize the active components of CM, aliquots were (i) heat-treated (95°C/5 min) and compared with freshly thawed CM that was not subjected to heat, (ii) treated with 8 μg (~10% of total protein) proteinase K (Zymo Research, Irvine, CA) versus vehicle (proteinase K storage solution), or (iii) centrifuged through 30-kDa molecular weight cutoff filters according to the manufacturer’s instructions (Microcon YM30; Millipore Corp.). Cutoff at 30 kDa was chosen because previous characterization of epithelia plus CM bacteria showed that some <20-kDa cytokines were found in the retentate instead of the expected flowthrough (63). This suggested that in nonnaturading conditions, some proteins aggregate prior to molecular weight cutoff. Filters were washed with sterile water, and the flowthroughs were pooled and concentrated to the original volume using vacuum centrifugation (SPD 1010 SpeedVac; Thermo Fisher). Following the filter washes, the desalted retentates (>30-kDa fraction) were collected and restored to the original volume with SNM/0.1% serum. These processed CMs were inoculated with *S. aureus* as described above, and *S. aureus* CFU were quantified after incubation at 37°C/5% CO2 for 24 h.

**Statistical analysis.** Data were analyzed using GraphPad Prism 8 software (GraphPad Software, La Jolla, CA). In most cases, *S. aureus* counts (CFU/treatment well) were expressed as percent recovery; *S. aureus* CFU yield for *S. aureus* alone (no competitor) were set at 100% and compared to *S. aureus* yield when admixed with test strains. In Fig. 4 and 6 to 8, data were analyzed by one-way analysis of variance (ANOVA). For Fig. 3, groups were compared using unpaired t tests (two-tailed). Correlations (Pearson r, R2, P value in Fig. 1) between nasal *S. aureus* counts and *S. aureus* growth *in vitro* were considered significant if P < 0.05, and linear regression line was significant for a nonzero slope.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**FIG S1**, PDF file, 0.1 MB.

**FIG S2**, PDF file, 0.1 MB.

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We declare no conflicts of interest.

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