The Human Nasal Microbiota and *Staphylococcus aureus* Carriage

Daniel N. Frank¹,², Leah M. Feazel¹, Mary T. Bessesen²,³,⁴, Connie S. Price⁴,⁵, Edward N. Janoff²,³,⁴, Norman R. Pace¹,²,⁶

¹Department of Molecular, Cellular, and Developmental Biology, University of Colorado at Boulder, Boulder, Colorado, United States of America, ²Mucosal and Vaccine Research Program Colorado, Aurora, Colorado, United States of America, ³Department of Veterans Affairs Medical Center, Denver, Colorado, United States of America, ⁴University of Colorado School of Medicine, Aurora, Colorado, United States of America, ⁵Division of Infectious Diseases, Denver Health Medical Center, Denver, Colorado, United States of America

Abstract

**Background:** Colonization of humans with *Staphylococcus aureus* is a critical prerequisite of subsequent clinical infection of the skin, blood, lung, heart and other deep tissues. *S. aureus* persistently or intermittently colonizes the nares of ~50% of healthy adults, whereas ~50% of the general population is rarely or never colonized by this pathogen. Because microbial consortia within the nasal cavity may be an important determinant of *S. aureus* colonization we determined the composition and dynamics of the nasal microbiota and correlated specific microorganisms with *S. aureus* colonization.

**Methodology/Principal Findings:** Nasal specimens were collected longitudinally from five healthy adults and a cross-section of hospitalized patients (26 *S. aureus* carriers and 16 non-carriers). Culture-independent analysis of 16S rRNA sequences revealed that the nasal microbiota of healthy subjects consists primarily of members of the phylum Actinobacteria (e.g., *Propionibacterium* spp. and *Corynebacterium* spp.), with proportionally less representation of other phyla, including Firmicutes (e.g., *Staphylococcus* spp.) and Proteobacteria (e.g., *Enterobacter* spp.). In contrast, inpatient nasal microbiotas were enriched in *S. aureus* or *Staphylococcus epidermidis* and diminished in several actinobacterial groups, most notably *Propionibacterium acetogenes*. Moreover, within the inpatient population *S. aureus* colonization was negatively correlated with the abundances of several microbial groups, including *S. epidermidis* (p = 0.004).

**Conclusions/Significance:** The nares environment is colonized by a temporally stable microbiota that is distinct from other regions of the integument. Negative association between *S. aureus*, *S. epidermidis*, and other groups suggests microbial competition during colonization of the nares, a finding that could be exploited to limit *S. aureus* colonization.

Introduction

*Staphylococcus aureus* is an invasive human pathogen with increasing incidence and morbidity in hospitals and the community. Both healthy persons and those with underlying illness are at risk for diverse skin and soft tissue infections, endocarditis, osteomyelitis, meningitis, bacteremia, and pneumonia (including pneumonia arising as a complication of influenza [1]), with mortality rates ranging from 6–40% [2,3]. The high frequency of poorly responsive and recurrent *S. aureus* disease in apparently immunocompetent hosts is a challenging feature of these infections [4]. Groups that are particularly susceptible include children in daycare [5], sports teams [6,7,8,9], jailed inmates [10,11,12], and military personnel [13,14,15,16]. Moreover, the emergence and rapid spread of methicillin-resistant *S. aureus* (MRSA) has placed substantial burden on the healthcare system.

Colonization of the nares is a potent and increasingly prevalent risk factor for subsequent *S. aureus* infection [17,18,19,20]. In at least 80% of *S. aureus* bacteremia cases in colonized subjects, the infecting strain is identical to a nasal colonizing strain detected prior to onset of bacteremia [17,21]. Followed longitudinally, approximately 20–30% of persons are colonized persistently with *S. aureus*, 30% are colonized intermittently, and 50% never, or rarely, are colonized [22,23]. Why some individuals apparently are resistant to colonization, and thus at lower risk of infection, remains an open question. Understanding the biology of this pathogen, especially its ecological niche in humans and the initial step in infection, colonization, may therefore provide new modalities to limit pathogenesis.

*S. aureus* carriage is influenced by myriad host and environmental factors [24,25]. To establish itself in the nares, *S. aureus* must successfully compete with many co-occurring microorgan-
isms, including corynebacteria, coagulase-negative staphylococci, and Streptococcus pneumoniae [26]. We hypothesize that competition and cooperation between S. aureus and nares-associated microbial communities directly impacts the incidence and prevalence of S. aureus colonization and subsequent infection. Although prior studies have analyzed associations between S. aureus and other well-characterized microorganisms [27,28,29], the microbial consortia that normally inhabit the nasal cavity may be more complex than indicated by traditional microbiological culture. Consequently, the microbial ecology of S. aureus colonization likely is incompletely understood. To surmount these potential limitations, we used culture-independent analyses of 16S ribosomal RNA sequences to more fully characterize the repertoire of indigenous microbial communities within the human nares of healthy and hospitalized adults in relation to S. aureus colonization.

Results

Study Design

To determine the frequency, diversity, and temporal stability of resident microbial communities, we collected nasal specimens longitudinally from the left and right nares of five healthy adults (Subjects A–E) over the course of 2–24 weeks (timepoints are listed in Fig. 1). For comparison, axilla, groin, and nasal specimens were collected in parallel from one individual (Subject A) to assess whether the nares harbor the same types of microorganisms as other regions of the integument. All healthy adults showered daily, 4/5 used antiperspirants and 1/5 used deodorant daily.

Hospitaled adults were sampled in a case-control study of 20 MRSA-colonized patients and 24 patients without MRSA colonization (based on routine nasal swab culture) while admitted to the intensive care units in two hospitals. Colonization was identified by results of culture of both nares with nasal swabs inoculated onto CHROMagar (Materials and Methods). Because patients without MRSA by culture could be either colonized with methicillin-sensitive S. aureus (MSSA) or not colonized by S. aureus, DNA prepared from nasal swabs was subjected to S. aureus-specific femA gene PCR [30] to detect all S. aureus (i.e., MSSA and MRSA). Two specimens were excluded on the basis of poor DNA recovery (i.e., negative rDNA PCR results, described below). All MRSA culture-positive patients were positive for S. aureus by femA PCR, indicating a low false negative rate (<1/16) of this PCR assay. Patients with negative cultures for MRSA and positive S. aureus PCR results were presumptively classified as MSSA. In aggregate, 18 inpatients were classified as MRSA carriers, 8 as MSSA carriers, and 16 S. aureus non-colonized.

Culture-Independent Microbe Identification

Bacteria present in specimens were identified by phylogenetic analysis of rDNA sequences, amplified from DNA isolated from swabs by PCR with pan-bacterial primers. Among healthy adults, PCR amplification was successful in 63/74 (85%) nasal specimens and 9/9 (100%) groin specimens, but only 4/11 (36%) axilla specimens. The failure of some samples to amplify was likely due to insufficient biomass. Although sampling methods could have been optimized to increase biomass (e.g., through wet-swabs or lavage), we followed the standard procedures used in the hospitals.
in order to compare data from all specimens. PCR successfully detected bacterial 16S rDNA in 26/28 (93%) inpatients colonized with *S. aureus* and 16/16 (100%) without *S. aureus* colonization.

Broad-range rRNA amplicons were subjected to high-throughput pyrosequencing on the Roche 454 GS-FLX platform. The reverse rRNA PCR primer (5'-3' SSU338R [31]) included a barcode sequence unique to each specimen in order to amplify and quantify multiple bacterial 16S rDNA sequences simultaneously (multiplex pyrosequencing reactions) [32]. To maximize the accuracy of bacterial identification, we rigorously screened sequences to remove poor quality reads. The resulting dataset consisted of 33,289 polished sequences, with a median length of 237 nucleotides (range 177–268 nt) and a mean of 268 sequences per specimen. Good’s coverage values ranged from 88%–100% per subject (mean 98%) therefore the true biodiversity of each specimen was adequately sampled.

rRNA sequences were provisionally identified by BLAST query of 16S rRNA sequences extracted from the All-Species Living Tree Project database (version LTP S95; [33]) and results corroborated through the RDP Classifier tool [34,35], parsimony insertion into the SILVA SSURef_98 tree [36], and pairwise distance comparisons to aligned staphylococcal sequences. Although the LTP database contains only a fraction of the 16S rRNA sequences present in GenBank or EMBL, its focus on type strains, rather than environmental sequences, can provide more accurate taxonomic assignments of human-associated bacterial rDNA, which typically are closely related to well-characterized microbial taxa [37,38,39]. Indeed, 91% of the rRNA sequences were assigned to the species-level (BLAST %ID scores ≥97%) and 93% assigned to the genus-level (BLAST %ID scores ≥95%). The remaining sequences were classified to higher taxonomic levels. Excellent concordance was observed between BLAST, RDP, and SILVA classifications (<0.1% of sequences had discordant results, which were resolved in favor of matches between two of the three methods).

**Ecology of the Healthy Nares Microbiota**

The majority of nares rRNA sequences obtained from healthy individuals belonged to only two bacterial phyla, the Actinobacteria (i.e., High-G+C Gram positive organisms such as corynebacteria; 68% of sequences) and Firmicutes (i.e., Low-G+C Gram positive organisms such as staphylococci; 27% of sequences; Table 1 and Table S1). Proteobacteria (4.1%), Bacteroidetes (1.4%), Fusobacteria (0.21%), Cyanotharios (0.08%), Tenericutes (0.07%) and Deinococcus (0.01%) accounted for the remainder of phylum-level diversity. Members of the Actinobacteria and Firmicutes also dominated specimens obtained from the groin and axilla (Fig. 1, Table 1, Table S1). Similar distributions of predominant phyla have been reported for integument-associated microorganisms of healthy individuals [37,40,41] and those with chronic wounds [39].

Although the precise distributions of species-level microbial groups differed from specimen to specimen, consistent patterns of microbial taxa were observed over time within each individual and anatomical location (Fig. 1, Fig. 2). For instance, all of the longitudinal groin specimens from Subject A were dominated by corynebacterial species (e.g. *C. mucifaciens, C. minutissimum*) that were not present, or much less abundant, in the axilla and nares samples of Subject A, including specimens collected on the same day. Similarly, sequences representative of *Propionibacterium acnes* were prevalent in the longitudinal nares samples of all subjects, but were observed far less frequently in axilla and groin swabs. *Staphylococcus epidermidis* was prevalent in the nares and axilla specimens, but not the groin, whereas *S. aureus* was observed only in the nares of two individuals. Specimens collected at the same time point from the left and right nares of the same individual did not differ appreciably in community composition relative to one another (Fig. 1), therefore these sequences were pooled for subsequent analyses.

Quantitative assessment of similarities between populations (using the Morisita-Horn similarity index) indicated that the nares, axilla and groin harbor distinct and temporally stable microbial communities (Figs. 2 and 3, Table 2). In general, nares samples were more similar to one another than to groin or axilla specimens, even when collected from the same host on the same day (e.g., p<0.001 for intra-subject comparisons of axilla vs. groin, axilla vs. nares, and groin vs. nares in Subject A; Table 2). Interestingly, the axilla and groin microbiotas of Subject A were no more similar to the nares microbiota of Subject A than to those of Subjects B-E (Fig. 2 and 3, Table 2).

The nares microbiotas of Subjects B, C, and D were indistinguishable from one another, whereas those of Subjects A and E were unique (Figs. 1, 2, and 3). For example, despite some temporal variability, the longitudinal nares samples of Subject A were significantly more similar to each other than to the nares microbiota of the other subjects (p<0.001 for within-subject to between-subject comparisons; Table 2). Of possible note, Subject A and E are male, Subjects B-D are female, and Subjects A and B and Subjects D and E are co-habitating couples. Thus, cohabitation did not result in convergence of nares microbiotas. Both Principal Components Analysis and hierarchical clustering corroborated these results (Fig. 4) and thereby provide additional evidence of anatomy-specific microbiotas.

The temporal variation observed in microbial populations could represent either microbial succession, as newly colonizing microbes displaced other species, or stochastic fluctuations about a fairly constant mean. The latter model, rather than succession, is more consistent with these data because communities remained generally similar to baseline samples over time (Fig. 5).

**Ecology of the Nares Microbiota in Hospitalized Patients**

Similar to healthy adults, most of the phylum-level nasal biodiversity observed in the inpatient population was restricted to the Firmicutes (71% of inpatient sequences) and Actinobacteria (20% of inpatient sequences). However, Firmicutes were significantly more abundant in inpatients compared to healthy adults (71% vs. 27% sequence abundance, respectively; Table 1 and Table S2), whereas Actinobacteria were concomitantly less abundant (20% vs. 68% for inpatients vs. healthy).

Differences in the abundances of Firmicutes between inpatients and healthy adults (Table 1) were due primarily to increased abundances of just two species, *S. aureus* and *S. epidermidis*, which together accounted for ≥50% (10,201/18,080) of the inpatient sequences. Similarly, most of the reduction in Actinobacteria in inpatients was associated with significantly diminished abundance of *Propionibacterium acnes* (Table 1).

No differences were apparent between the nasal microorganisms of MRSA and MSSA colonized individuals (data not shown), therefore datasets were combined to compare with non-colonized individuals. Among inpatients, *S. aureus* colonization was negatively correlated with *S. epidermidis* abundance (p = 0.004; Table 1). *S. aureus* sequences were the most abundant sequence-types encountered in patients classified as *S. aureus*-carriers by femA PCR, whereas *S. epidermidis* sequences were most abundant in the non-carriers (Table 1, Fig. 6). These results both confirm the diagnostic utility of the femA PCR assay used to classify patients and indicate that both staphylococcal species can dominate the nares microbiota of certain inpatients. Although several other
Table 1. Nares-associated bacterial diversity in healthy and hospitalized adults 1.

| Inpatient | S. aureus Carriage 2 |
|-----------|----------------------|
| **Top Blast Hit 3** | Neg | Pos | ρ 4 | Healthy |
| **Firmicutes** | 54.0% 5 | 66.5% 5 | < 0.001 | 25.6% 5 |
| Staphylococcus aureus | 0.1 | 46.1 | 4.5 |
| Staphylococcus epidermidis | 43.7 | 8.2 | 0.004 | 10.3 |
| Peptostreptococcus spp. | 1.8 | 1.3 | 3.2 |
| Anaerococcus spp. | 1.1 | 1.0 | 1.4 |
| Other Firmicutes | 7.3 | 9.8 | 6.2 |
| **Actinobacteria** | 34.4 | 14.9 | 0.06 | 69.0 |
| Propionibacterium acnes | 12.2 | 3.4 | 42.4 |
| Corynebacterium accolens | 5.6 | 5.2 | 7.3 |
| Corynebacterium pseudodiphtheriticum | 0.0 | 1.9 | 4.8 |
| Corynebacterium tuberculostearicum | 5.7 | 1.0 | 8.0 |
| Mycobacterium spp. | 0.1 | 0.2 | 3.4 |
| Other Actinobacteria | 10.8 | 3.1 | 3.1 |
| **Proteobacteria** | 8.6 | 17.0 | 4.0 |
| Enterobacter ludwigii | 0.1 | 5.3 | 0.1 |
| Other Proteobacteria | 8.5 | 11.7 | 3.9 |
| **Bacteroidetes** | 1.11 | 0.34 | 1.36 |
| **Fusobacteria** | 0.08 | 0.02 | 0.21 |
| **Cyanobacteria** | 0.02 | 0.02 | 0.08 |
| **Deinococcus** | 0.02 | 0.00 | 0.01 |
| **Tenericutes** | 0.04 | 0.00 | 0.07 |
| Sequences 6: | 5227 | 9095 | 16411 |
| Subjects 7: | 16 | 26 | 5 |
| Specimens 8: | 16 | 26 | 63 |
| Observed 99% OTUs 9: | 36.8 | 34.7 | 52.4 |
| Estimated 99% OTU richness 10: | 108.0 | 113.8 | 244.6 |
| Shannon diversity: | 3.5 | 3.4 | 4.5 |
| Shannon evenness: | 68% | 67% | 79% |
| Simpson diversity: | 6.6 | 6.9 | 14.1 |

1Table summarizes most abundant species/genera of 16S rRNA sequences in anterior nares swabs. See supplemental information for complete dataset.
2S. aureus non-colonized (Neg) with colonized (Pos) ICU patients, classified by culture and femA gene PCR.
3Inferred from highest bit-score in BLAST query. Blast %IDs < 97 are named only to the genus level. Sequence abundances are listed for phyla (bold) and the 10 most abundant species/genera (italics; 90% of total sequences).
4p-value for Student’s t-test comparison of colonized and non-colonized patients. Only values < 0.1 are noted.
5Abundance of sequences for each category. Values are averages for subjects in a category.
6Number of sequences analyzed in each category.
7Subjects included in category.
8Specimens included in category. If more than one specimen/subject were analyzed, mean rRNA abundances were weighted by the number of specimens so that all subjects contribute equally to the mean value.
9Operational taxonomic units (OTUs) were assembled by complete-linkage clustering at a threshold of 99% uncorrected sequence identity. Tabulated values are means obtained through 1000 boot-strap replicates using rarefaction to normalize for differences in sampling efficiency.
10Schao1 non-parametric estimate of species richness for 99% OTUs.

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species, such as Corynebacterium spp., also were less abundant in S. aureus-colonized subjects, this study was not sufficiently powered to gain statistical significance. However, the phyllum Actinobacteria as a whole was reduced in abundance in S. aureus colonized inpatients (p = 0.06).

The dataset presented in Table 1, which pools data from many subjects, masks substantial individual-to-individual variation in nares populations (Fig. 6). The negative correlation between the abundances of S. aureus and S. epidermidis in S. aureus colonized and non-colonized inpatients is readily apparent (Fig. 6A), as is the significant loss of Actinobacteria in inpatients compared with healthy adults (Fig. 6B). In only one patient (V11) were both S. aureus and S. epidermidis present in relative abundances > 25%. Furthermore, a subset of inpatients exhibited elevated levels of Proteobacteria (mainly members of the orders Enterobacteriales and Pseudomonadales) and Firmicutes other than S. aureus and S. epidermidis. Thus, blooms in S. aureus or S. epidermidis were not characteristic of all inpatients, as might be expected given the
range of morbidities that likely were exhibited by the inpatient population.

In a zero-sum assay such as broad-range PCR, a bloom of one type of microbe would lower the proportions of all other microbes in a sequence library, regardless of whether the bloom actually affected the growth of the other microbes. However, removal of \textit{S. aureus} sequences from the sequence datasets revealed different distributions of microbial groups in \textit{S. aureus} colonized compared with non-colonized individuals (Fig. 7). Of most relevance, the abundance of \textit{S. epidermidis} was greatly reduced relative to most other bacterial taxa in \textit{S. aureus} colonized individuals. This suggests that \textit{S. aureus} may alter the composition of the underlying nares bacterial communities, rather than simply grow without impacting or displacing other microbial communities.

**Reduced Biodiversity in Inpatient Nares Microbiota**

Healthy adults harbored significantly more species-rich and diverse nares microbiotas than did hospitalized individuals (Table 1, Fig. 8). Approximately twice as many species-level microbial groups (defined as 99\% OTUs) were identified in healthy adult nares microbiotas as were found in inpatient microbiotas (p<0.001; 245 vs. 112 OTUs, respectively; Table 1) as estimated by the \textit{S$_{chao1}$} non-parametric species-richness index (p<0.001; \textit{S$_{obs}$}: 52 vs. 36 OTUs, respectively; Table 1, Fig. 8). Comparison of Shannon diversity estimates (Table 1, Fig. 8) indicated that inpatient nares microbiotas also were both less diverse (p<0.05) and less even (p<0.05) than healthy microbiotas. Each of these results is consistent with the observed blooms in \textit{S. aureus} and \textit{S. epidermidis}.

**Accuracy of Sequence-based Classification of \textit{Staphylococcus} Species**

As described above, sequences were classified and clustered on the basis of BLAST results. Similar results also were obtained when sequences were clustered into 97\% and 99\% operational taxonomic units (OTUs). We chose to present BLAST-based
comparisons, because they tend to mitigate possible overestimation of OTU-based clusters that can result from sequencing errors, ambiguous alignment, and/or intra-species chimeras [42, 43]. Because the strength of our conclusions with respect to correlations between the abundances of *S. aureus* and other nares-associated species (particularly *S. epidermidis*) is dependent on the accuracy of the rDNA sequence-based microbe identification, we sought to corroborate the results of *S. aureus* and *S. epidermidis* classifications through more detailed sequence analysis.

Because relatively short (<250 nt) pyrosequencing reads might not be as phylogenetically informative as longer sequences, we cloned and Sanger sequenced broad-range amplicons generated from each inpatient sample using the primers 8F and 805R [31], which samples approximately one-half of the 16S rRNA sequence. A total of 3758 Sanger sequences were generated and assigned to taxa in parallel with the pyrosequencing reads discussed above. In general, similar microbial distributions were observed between the two sequencing platforms, despite the use of different PCR primer sets and phylogenetic analysis of partially overlapping sections of the 16S rRNA sequence (Table S2). Indeed, the relative abundances of most taxa varied by less than five-fold between platforms. Of note, *S. aureus* and *S. epidermidis* abundances were qualitatively similar, and thus indicated no systematic bias due to shorter pyrosequencing reads. However, several exceptions were apparent, most notably *Propionibacterium acnes*, which was the 3rd most abundant group in the pyrosequencing dataset, but present at much reduced levels in the Sanger dataset. These differences may have resulted from primer bias or sequence-length dependent ambiguities in classification.

To further corroborate taxonomic assignments, each presumptive *S. aureus* and *S. epidermidis* sequence was aligned to the SILVA SSURef database (v. 98), then pairwise similarities were calculated against staphylococcal sequences in the Living Tree Project (LTP_S95) rDNA database. For each nares sequence, its highest similarity was noted in comparison to 1) the subset of cognate sequences in the LTP_S95 database (i.e., presumptive *S. aureus* sequences in our dataset vs. *S. aureus* sequences in the database) and 2) all other staphylococcal sequences in the LTP_S95 dataset.

Figure 9 presents scatterplots of these two similarity scores plotted for each nares sequence (Panel A for *S. aureus* and Panel B for *S. epidermidis*). In all instances, the nares sequences were most similar to sequences from the predicted species, despite the variability in raw similarity scores. Because the genomic 16S rRNA gene sequences of *S. aureus* and *S. epidermidis* differ by only ca. 1.5%, we were surprised that nares sequences with BLAST %ID or pairwise similarity scores less than 98.5% were most closely related to *S. aureus* or *S. epidermidis*. We attribute this seeming discrepancy to pyrosequencing errors [42, 43] and inaccuracies in sequence alignment, both of which would generally depress all pairwise similarity scores, yet likely preserve the identity of the true nearest neighbor sequence in this type of analysis.

### Discussion

Determination of the biological factors that naturally protect individuals from *S. aureus* colonization may lead to novel strategies for preventing infections. As a first step towards understanding the microbial ecology of *S. aureus* carriage, we have analyzed the nares microbiotas in cohorts of healthy and hospitalized individuals. Patients were classified as *S. aureus* colonized cases or *S. aureus* non-colonized controls, from which we determined and correlated the point-prevalences of *S. aureus* and other microbes in the nares. The nares microbiotas of a small cohort of healthy adults were tracked longitudinally to determine the types and temporal variability of microorganisms that normally inhabit the anterior nares. Although this healthy cohort was not designed to be a control for the inpatient study, it nevertheless provides important context in which to interpret the results of the inpatient point-prevalence survey. For instance, it was critical to first establish whether the anterior nares are inhabited either by a defined, characteristic

| Table 2. Community similarity (Morisita-Horn Index) comparisons for healthy adults.* |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Subject A | Axilla | Groin | Nares | Nares | Nares | Nares | Nares |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Subject A | 0.87 | 0.11 *** | 0.13 *** | 0.14 *** | 0.34 ** | 0.27 *** | 0.26 *** |
| Subject B | 0.11 *** | 0.78 | 0.03 *** | 0.01 *** | 0.06 *** | 0.03 *** | 0.05 *** |
| Subject C | 0.13 *** | 0.03 *** | 0.67 | 0.31 *** | 0.36 *** | 0.46 *** | 0.40 *** |
| Subject D | 0.14 *** | 0.01 *** | 0.31 *** | 0.95 | 0.91 | 0.79 *** | 0.54 |
| Subject E | 0.34 *** | 0.06 *** | 0.36 *** | 0.91 | 0.97 | 0.84 * | 0.61 |
| Subject A | 0.27 *** | 0.03 *** | 0.46 *** | 0.79 *** | 0.84 ** | 0.92 | 0.63 |
| Subject E | 0.26 *** | 0.05 *** | 0.40 *** | 0.54 *** | 0.61 * | 0.63 *** | 0.66 |

*Median Morisita-Horn scores for all pairwise combinations of specimen types. Statistical significance assesses whether within-group similarity scores (i.e., axilla specimens vs. axilla specimens), which are in bold, differ significantly from without-group mean similarity scores in the same column (i.e., axilla-specimens vs. groin specimens), measured by Wilcoxon rank-sum test. In this case, the axilla-axilla score of 0.83 differs significantly from the axilla-groin score of 0.14. Significance levels are indicated by ***p<0.001; **p<0.01; *p<0.05; p<0.05. doi:10.1371/journal.pone.0010598.t002
microbiota or by transient microbial assemblages that result from perpetually changing exposures to bioaerosols, dust particles, water, etc.

Few in-depth longitudinal studies of the human microbiota have been reported to date [44]. Our results indicate that the dominant microorganisms of the axilla, groin, and anterior nares remain relatively constant on the time-scale of weeks to months (Figs. 1 and 2), in the absence of mitigating factors. Although each individual was host to different suites of microorganisms, many commonalities were observed between

Figure 4. Similarity of microbiotas determined longitudinally for healthy adults. Panel A. Principal Components Analysis of Microbiotas. Colors indicate the subject and anatomical location from which longitudinal specimens were obtained. No sampling time-dependent trends were observed in the data, so datapoints are not labeled with respect to time of collection. Panel B. Hierarchical Clustering. Colors indicate the subject and anatomical location from which longitudinal specimens were obtained. Leafs are labeled by subject and day of collection. See Materials and Methods for details. doi:10.1371/journal.pone.0010598.g004
individuals, such as nasal carriage of \( P. \) acnes, \( S. \) epidermidis, and corynebacterial species. Collectively these microorganisms define a pan-microbiota of the human anterior nares as has been reported for other anatomical sites, including the genitourinary tract [43], gut [38,46,47,48], lung [49], and skin [37,40,50,51,52].

The nares microbiotas of most hospitalized patients differed substantially from those of the healthy cohort in the kinds and diversities of prevalent microbes. To our knowledge, this is the first characterization of the human nasal microbiota in a clinical context. Our results broadly define three types of microbial populations within the nasal cavity (Fig. 10). First, healthy adults and a subset of inpatients harbored nares communities dominated by Actinobacteria (mainly Propionibacterium and \( \)Corynebacterium spp.), with fewer staphylococci. Second, in the majority of \( S. \) aureus-colonized inpatients \( S. \) aureus was the dominant nasal species, with concomitant reductions in the prevalences of Actinobacteria. Third, many \( S. \) aureus non-colonized patients carried \( S. \) epidermidis as the dominant species, accompanied by reduced levels of Actinobacteria. Thus in this study, \( S. \) aureus carriage was negatively associated with a variety of other nares-associated microbial species, most significantly \( S. \) epidermidis and \( P. \) acnes (Table 1). These results are consistent with published reports that selected organisms can interfere with \( S. \) aureus colonization [26,29]. However, the culture-independent, high-throughput strategy utilized in this study permitted detailed characterization of whole populations of nasal microorganisms, rather than individual species, in relation to \( S. \) aureus occurrence.

We propose that the underlying morbidities of the inpatient population and/or their exposure to the hospital environment caused a transition from a “healthy” nasal microbiota to either an \( S. \) aureus- or \( S. \) epidermidis-dominated microbiota (Fig. 10). Development of an \( S. \) epidermidis-dominated microbiota may either protect a person from subsequent \( S. \) aureus colonization or be an intermediate stage in succession to a nasal microbiota dominated by \( S. \) aureus. In the latter model, loss of commensal species, such as \( P. \) acnes, rather than gain of \( S. \) epidermidis could be the key determinant of subsequent \( S. \) aureus colonization. Because only point-prevalences of microbial groups were determined for inpatients in this study, we could not ascertain whether particular organisms interfere with or promote \( S. \) aureus colonization; longitudinal studies will shed additional light on the potential for particular microorganisms to inhibit acquisition of \( S. \) aureus (or vice versa). As competing species and strains are identified, the molecular and physiological factors that influence adaptation and competition within the nares environment can be studied in greater depth. For instance, selection for antibiotic resistance and exchange of resistance-encoding genes are expected to be critical factors in nares ecology in the clinical context [53,54].

The inpatient study population, which was drawn from ICUs, undoubtedly is confounded by myriad co-morbidities, therefore follow-up studies also are necessary to delineate the interplay between specific clinical factors, the nares microbiota, and the risk of \( S. \) aureus colonization. Nevertheless, similar findings were obtained at two medical centers, suggesting that the results were not unique to a particular patient population or due to methodological bias. Rather, the underlying morbidities of the inpatient study population or exposure to the hospital environment likely account for differences observed between healthy and inpatient cohorts.

Because colonization by \( S. \) aureus is a significant predisposing factor for subsequent infection, deeper understanding both of colonization and the ecological niche from which infections originate (the nares) could lead to novel prevention strategies. Statistically significant enrichment of particular microbial groups (e.g., \( S. \) epidermidis) in individuals who are not colonized by \( S. \) aureus suggests that specific bacterial products interfere with colonization, either by competition for local resources, by antibacterial activity, or by competitive attachment to mucosal sites. Definitive identification and characterization of interfering bacteria or their products may provide candidates for novel medical interventions, both passive and active (e.g., as vaccines or probiotics). For instance, protective microorganisms might be developed for use as probiotic agents within the nasal cavity. Alternatively, the products...
A. S. aureus and S. epidermidis

B. Other predominant bacterial groups

Figure 6. Inter-subject variability in nares microbiota. Panel A. Comparison of S. aureus and S. epidermidis proportions in rDNA libraries. Column heights represent relative abundances of particular microbial groups in hospitalized patients and healthy adults. Hospitalized patients are labeled “D” or “V”, based on the hospital ICU at which they were admitted. The dashed lines denote the mean abundances of S. aureus and S. epidermidis -- 4.8% and 9.3% respectively -- in all nares specimens from healthy adults. rDNA sequences were determined by pyrosequencing. Panel B. All other bacteria, grouped at taxonomic-order level. Dashed line represents the mean abundance of Actinomycetales in all nares specimens from healthy adults.

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of S. aureus-interfering genes could have potential as topical anti-S. aureus agents that block essential binding sites or generate local innate immune activity. The targets of specific inhibitory factors might also suggest candidate antigens for vaccines to prevent S. aureus colonization and therefore mitigate the effects of this increasingly prevalent and virulent pathogen.

Materials and Methods

Human subjects and sample collection

The study was approved by the Institutional Review Board of the University of Colorado, Boulder (protocols 0807.21 and 1009.35). Prospectively sampled healthy adults provided written informed consent.
consent to be enrolled in the study. Specimens collected from hospitalized patients were deidentified of all personal information, so informed consent was not required. Consequently, we were unable to collect clinical or demographic information about the patients beyond MRSA culture results. Nasal swabs were collected through convenience sampling of patients admitted to the intensive care units of two academic medical center hospitals in the Rocky Mountain U.S. region that routinely survey inpatients for nasal MRSA carriage. Specimens were obtained with sterile, dry swabs (Becton, Dickinson and Company), which were rotated five times around the inside of both nostrils while applying constant pressure. Specimens were randomly selected from those collected at the time of admission to intensive care units or ward nursing units, at the time of inter-unit transfer, at time of discharge, or on hospital days 7, 14, 21 etc. Inpatient nasal swabs were used to inoculate MRSA culture plates then promptly preserved in 70% ethanol and stored at –20°C. Nasal swabs from healthy subjects were self-administered in a similar manner, however, right and left nares were sampled separately. Specimens from the axilla and groin were collected by dry swabbing for 5–10 seconds. All swabs were frozen (–20°C) upon collection from healthy individuals.

**MRSA culture**

Nasal swabs were cultured on CHROMagar plates (Becton, Dickinson and Company) in a non-CO₂ incubator and deemed positive if mauve-colored colonies were detected after 18–48 hours incubation. *S. aureus* was confirmed by Gram stain, catalase assay, and/or latex agglutination test.

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**Figure 7. Impact of *S. aureus* sequences on distributions of other nares bacteria.** Panel A. Percent abundance of top twelve genera. Panel B. Percent abundance of top twelve genera adjusted by removal of *S. aureus* from total sequence counts.

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**Figure 8. Ecological richness and diversity of nares microbiotas.** The top panel presents the distributions of species richness indices ($S_{obs}$) calculated for nares specimens obtained from hospitalized or healthy adults. The middle panel summarizes sample biodiversity (Shannon diversity index $H_o$) and the lower panel presents species evenness ($H_o/H_{max}$). $H_o$ = Shannon index; $H_{max}$ = maximum value of $H_o$ for a specimen). Significance levels are indicated by * $p<0.05$; ** * $p<0.001$.

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Figure 9. Accuracy of rDNA sequence-based classification of S. aureus and S. epidermidis. Each nares sequence classified as S. aureus or S. epidermidis by BLAST query of a highly-curated database (SILVA LTP_S95) was aligned and its pairwise sequence similarity determined in relation to each sequence in a dataset consisting of all LTP_S95 staphylococcal sequences along with other staphylococcal genomic sequences. Maximum percent identity scores were identified both to the predicted species (S. aureus or S. epidermidis) and to all other staphylococci in the dataset. These two scores were plotted for each nares sequence. Points falling above the diagonal line therefore represent nares sequences that were most closely related to the species predicted by BLAST, whereas points falling below the diagonal suggest mis-classification. The majority of sequences were at least 97% identical to the predicted species, which justifies species-level identification of rRNA sequences. We interpret the low maximum percent identity values obtained for some sequences (i.e., 97% identity to any staphylococcal species, yet a top BLAST hit of S. aureus or S. epidermidis) as arising from base-calling or alignment errors inherent in analysis of pyrosequencing reads. Panel A. Nares sequences classified as S. aureus. Panel B. Nares sequences classified as S. epidermidis. doi:10.1371/journal.pone.0010598.g009

Figure 10. Distinct microbial populations in healthy and hospitalized adults. Pie charts depict average frequencies of dominant microorganisms in the anterior nares of healthy adults and inpatients, classified by S. aureus carriage status. Arrows outline possible pathways by which microbial populations develop in hospitalized patients. “Other” represents less abundant taxa, such as Proteobacteria and Firmicutes other than S. aureus and S. epidermidis. doi:10.1371/journal.pone.0010598.g010
DNA extraction

Swab heads were placed in sterile two-ml microcentrifuge tubes with 500 µl of TEN buffer (10 mM Tris-Cl pH 8.0, 1 mM EDTA, 1% NP40 nonionic detergent) and ca. 250 mg of zirconium beads (0.1 mm, Biospec Products Inc, Bartlesville, OK). Specimens were agitated in a Mini Beadbeater-8 (Biospec Products Inc, Bartlesville, OK) on the highest setting for 3 minutes, incubated at 95°C for 10 minutes, vortexed for 30 seconds, and then heated and vortexed a second time. The tubes were either placed on ice or stored at −20°C before PCR processing. Our previous studies indicate that this DNA extraction protocol is robust and efficiently lyses a variety of cell types, including Gram-positive bacteria, mycobacteria, and yeasts [37,39]. All DNA extraction and PCR steps were performed in a laminar flow hood that was decontaminated by UV light.

Multiplexed Pyrosequencing

30 µl PCR reactions contained 12 µl 2.5x HotMasterMix (5 PRIME Inc., MD, USA), 0.4 µM 27F-YM3 [35], 0.2 µM barcoded SSU338R [31] and 1 µl mixed-community genomic DNA. A single master PCR cocktail containing all reagents other than the genomic DNA and barcoded SSU338R [32] primers was set up then aliquoted into a 96-well PCR microtitre plate. PCR reactions were performed in duplicate using a protocol of 92°C 15 secs, 52°C 15 secs, and 65°C 45 secs. All samples were initially amplified through 30 cycles and additional cycles were performed if necessary for subsequent sequencing (38 cycles as the maximum).

PCR products were normalized prior to pyrosequencing with the SeqCapPrepTM Normalization Plate Kit (Invitrogen Inc., CA, USA), following the manufacturer’s protocol. Duplicate PCR reactions were pooled prior to normalization. Amplicons were eluted from normalization plates in 30 µl of 10 mM Tris-Cl (pH 8.0). 25 µl aliquots of each amplicon were combined in a pool, which was lyophilized to ~30 µl. One-half of the pool was electrophoresed through a 1.5% agarose gel in Tris/Acetate/EDTA and product excised with a sterile razor blade under low-wavelength ultraviolet light. DNA was eluted from the gel slice using the Montage™ DNA Gel Extraction Kit (Millipore Corp., MA, USA). This pooled DNA was provided to the Colorado Consortium for Comparative Genomics for pyrosequencing on a 454 Life Sciences GS-FLX instrument.

Barcoded, raw pyrosequencing reads were polished (bases with Q score<20 averaged across window of 5 nts. removed) and deconvoluted using the program bartab [32]. 32,341 sequences were excluded from analysis base on the following criteria: 1) trimmed length<175 nt; 2) >0 ambiguous base(s); and/or 3) absence of barcode or forward primer sequence. One specimen, D11, produced only 13 high-quality sequences and was removed from statistical analyses.

rDNA clone library construction and Sanger sequencing

SSU rRNA genes were amplified from DNA samples by PCR with primers specific for all bacterial SSU rRNA genes: 8F (5’AGAGTTTTGATCCTGCGCTCAG) and 805R (5’GACTACCAGGTTATCTTAT). Each 30 µl PCR reaction contained 12 µl 2.5x HotMasterMix (5 PRIME Inc., MD, USA), 25 ng of each primer, and 1 µl genomic DNA lysate. rDNA genes were amplified through thirty PCR cycles (92°C 30 sec., 52°C 60 sec., 72°C 90 sec.) and the expected products were inspected using ethidium-bromide-stained agarose gels (Kodak Inc.). PCR controls for each set of samples included extracts from unused, sterile swabs and sterile H2O (negative controls). PCR amplified DNA fragments were excised from agarose gels (1.5% agarose gel in tris-borate EDTA), purified using the QIAquick® gel extraction kit (Qiagen Inc., Valencia, CA), and cloned into the pCR4®-TOP® vector (TOPO® TA Cloning kit, Invitrogen Corp., Carlsbad, CA). For each clone library, 96 transformants were grown overnight at 37°C in 1.5 ml 2xYT medium using 96-well culture plates. To sequence the inserts of transformants, 20 µl of each overnight culture was mixed with 20 µl of 10 mM Tris-Cl (pH 8.0), heated 10 minutes at 95°C, and centrifuged 10 minutes at 1,360 x g in a 96-well plate centrifuge (Eppendorf Inc., Westbury, NY). One microliter of culture supernatant was used as template with vector-specific T7 and T3 primers in a 30 µl PCR reaction (38 cycles as above). Ten microliter of each PCR product were treated with the ExoSap-IT kit (USB Corp, Cleveland, OH) and cycle-sequenced using vector-specific T7 and T3 primers with the Big-Dye Terminator kit (Applied Biosystems, Inc., Foster City, CA). Sequencing was performed in-house on a MegaBACE 1000 (Amersham Biosciences, Piscataway, NJ) automated DNA sequencer. Sequence base calling and contig assembly were performed with the applications phred and phrap [56,57], as implemented by XplorSeq [58]. Vector and primer sequences were removed along with poor quality flanking nucleotides (Q<20).

Sequence Analysis

All sequences were aligned to the SILVA SSURef version 98 database using the SINA automated aligner provided by the SILVA web service [36]. Sequences with SILVA quality scores less than 75, including potential chimeric sequences, were removed from the dataset and not subjected to further analysis. Aligned sequences were inserted into the SSURef SILVA guide tree using the parsimony insertion function of ARB [59]. Initial taxonomic assignment of SSU sequences was made by a batch BLAST search of both GenBank and a local database of rRNA sequences extracted from the All-species Living Tree Project database (version LTP_S95; [33]) using the client applications blastcl3 and blastall (NCBI). Taxonomic lineage information was extracted from GenBank records of top BLAST hits and compared to results obtained from the Naïve Bayesian Classifier tool provided by the Riboosomal Database Project [34]. Ambiguities between GenBank and RDP classifications (<0.1% of sequences) were resolved by reference to the results of parsimony insertion into the SILVA guide tree in ARB.

To calculate ecological biodiversity indices, we assigned sequences to OTUs by nearest-neighbor clustering of uncorrected pairwise distances, using the application sortx and a distance cutoff of 99% [58]. The completeness of sequencing was assessed by Good’s Coverage estimator [60]. The evenness of sequencing was assessed by the rarefaction index [61]. The Chao1 diversity [62] was calculated using the program biodiv (D.N. Frank, unpublished. http://www.phyloware.com). Sequence distances between presumptive S. aureus and S. epidermidis 16S rRNA sequences and databased staphylococcal sequences were calculated and compared using the program xscmpdst (D.N. Frank, unpublished. http://www.phyloware.com).

Statistical analyses

All analyses used the R-software package [v.2.0.1; [63]]. Differences in the relative abundances of microbial groups between sample types were assessed by the two-tailed Student’s t-test for unequal variance (logit-transformation of abundance data did not alter the reported results for untransformed data). Morisita-Horn community similarity indices (G MH) were calculat-
ed using the vegdist function of the R package "vegan" [66].
Differences in C_MH values between categories were tested by Wilcoxon rank-sum test. Hierarchical clustering relied on the R package "cluster" [65] and were generated using both complete and average linkage. Dissimilarity matrices used as input were calculated using either the function "euclidean" (methods "euclidean" and "manhattan" distances) or vegdist (methods "jaccard" and "horn"). Heatmaps used the heatmap.2 function available through the R package "ggplot" [66].

DNA sequence accession numbers
Sequences were deposited in GenBank and assigned the accession numbers HM073607 - HM077276 and HM081990 - HM099519.

Supporting Information
Table S1

| References |
|-----------------------------------|
| 1. Kleves RM, Morrison MA, Nadler J, Petri S, Gerdman K, et al. (2007) Invasive methicillin-resistant Staphylococcus aureus infections in the United States. JAMA 298: 1763-1771. |
| 2. Nook GA, Rubin RJ, Schentag JJ, Klyuytman J, Hellblom EC, et al. (2007) National trends in Staphylococcus aureus infection rates: impact on economic burden and mortality over a 6-year period (1998-2003). Clin Infect Dis 45: 1132-1140. |
| 3. Nickerson EK, West TE, Day NP, Peacock SJ (2009) Staphylococcus aureus disease and drug resistance in resource-limited countries in south and east Asia. Lancet Infect Dis 9: 130-135. |
| 4. Kreisle K, Boyd K, Langenburg P, Rognham MC (2006) Risk factors for recurrence in patients with Staphylococcus aureus infections complicated by bacteraemia. Diagnostic Microbiology & Infectious Disease 53(3): 179-84. |
| 5. Velazquez-Guadarrama N, Martinez-Aguilar G, Galindo JA, Zuñiga G, Arbo-Sosa A (2009) Methicillin-resistant S. aureus colonization in Mexican children attending day care centres. Clin Invest Med 32: E57-63. |
| 6. Stacey AR, Endersby KE, Chan FC, Marples RR (1999) An outbreak of methicillin-resistant Staphylococcus aureus infection in a rugby football team. Br J Sports Med 32: 153–154. |
| 7. Nguyen DM, Maciia L, Brazzoli E (2005) Recurring methicillin-resistant Staphylococcus aureus infections in a football team. Emerg Infect Dis 11: 526-532. |
| 8. Kazaakova SV, Hagerman JC, Matava M, Shrivasaa A, Pielen L, et al. (2005) A clone of methicillin-resistant Staphylococcus aureus among professional football players. N Engl J Med 352: 468-475. |
| 9. Huijgens PW, van Lier AM, van Kregten E, Verhoef L, van Santen-Kersten M, et al. (2006) Methicillin-resistant Staphylococcus aureus in Dutch soccer team. Emerg Infect Dis 12: 1584-1586. |
| 10. Pan ES, Deep BA, Carleton HA, Charlebois ED, Sensabaugh GF, et al. (2003) Increasing prevalence of methicillin-resistant Staphylococcus aureus infection in Canadian children. JAMA 290(6): 716-20. |
| 11. Gilbert M, MacDonald J, Gregson D, Sihanshian J, Zhang K, et al. (2006) Outbreak in Alberta of community-acquired (USA300) methicillin-resistant Staphylococcus aureus in people with a history of drug use, homelessness or incarceration: CMAJ 175: 149-154. |
| 12. Main CL, Jayaratne P, Haley A, Rutherford C, Smaill F, et al. (2005) Outbreaks of infection caused by community-acquired methicillin-resistant Staphylococcus aureus in a Canadian correctional facility. Can J Infect Dis Med Microbiol 16: 343-348. |
| 13. LaMar JE, Carr RB, Zinderman C, McDonald K (2005) Sentinel cases of community-acquired methicillin-resistant Staphylococcus aureus onboard a naval ship. Mil Med 168: 139-140. |
| 14. Zinderman CE, Conner B, Malakooti MA, LaMar JE, Armstrong A, et al. (2004) Community-acquired methicillin-resistant Staphylococcus aureus among military recruits. Emerg Infect Dis 10: 941-944. |
| 15. Bradham GM, Schindler G, Skaeda D, Jensen B, McAllister S, et al. (2003) Emerging infections with community-associated methicillin-resistant Staphylococcus aureus in outpatients at an Army Community Hospital. Surg Infect (Larchmt) 6: 87-92. |
| 16. Ellis AW, Hospenthal DR, Dooley DP, Gray PJ, Murray CK (2004) Natural history of community-acquired methicillin-resistant Staphylococcus aureus colonization and infection in soldiers. Clin Infect Dis 39: 971-979. |
| 17. Wertheim H, Vos MC, Ott A, van Belkum A, Voss A, et al. (2004) Risk and outcome of nosocomial Staphylococcus aureus bacteremia in nasal carriers versus non-carriers. Lancet 364: 703-705. |
| 18. Perl TM, Callen J, Wenzel RP, Zimmerman MB, Pfeller MA, et al. (2002) Intranasal mupirocin to prevent postoperative Staphylococcus aureus infections. N Engl J Med 346: 1871-1877. |

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Table S2

| Author Contributions |
|-----------------------|
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Conceived and designed the experiments: DF ENJ NP. Performed the experiments: DF LMF. Analyzed the data: DF LMF ENJ NP. Contributed reagents/materials/analysis tools: DF MTB CSP. Wrote the paper: DF ENJ NP.
38. Frank DN, St Amand AL, Feldman RA, Boedeker EC, Harpaz N, et al. (2007) Molecular-phylogenetic characterization of microbial community imbalances in human inflammatory bowel diseases. Proc Natl Acad Sci U S A 104: 13780–13785.
39. Frank DN, Wysocki A, Specht-Glick DD, Rooney A, Feldman RA, et al. (2009) Microbial diversity in chronic open wounds. Wound Repair Regen 17: 163–172.
40. Gao Z, Tseng CH, Pei Z, Blaser MJ (2007) Molecular analysis of human forearm superficial skin bacterial biota. Proc Natl Acad Sci U S A 104: 2927–2932.
41. Grice EA, Kong HH, Conlan S, Deming CB, Davis J, et al. (2009) Topographical and temporal diversity of the human skin microbiome. Science 324: 1190–1192.
42. Kunin V, Engelbrektson A, Ochman H, Hugenholtz P (2010) Wrinkles in the rare biosphere: pyrosequencing errors can lead to artificial inflation of diversity estimates. Environ Microbiol 12: 118–123.
43. Quince C, Lanzen A, Curtis TP, Davenport RJ, Hall N, et al. (2009) Accurate determination of microbial diversity from 454 pyrosequencing data. Nat Methods 6: 639–644.
44. Dethlefsen L, Huse S, Sogin ML, Relman DA (2008) A diversity profile of the human skin microbiota. Genome Res 18: 1043–1050.
45. Paulino LC, Tseng CH, Strober BE, Blaser MJ (2006) Molecular analysis of fungal microbiota in samples from healthy human skin and psoriatic lesions. J Clin Microbiol 44: 2933–2941.
46. Aziz RK, Nizet V (2010) Pathogen microevolution in high resolution. Sci Transl Med 2: 16ps14.
47. Harris SR, Feil EJ, Holden MT, Quail MA, Nickerson EK, et al. (2010) Evolution of MRSA during hospital transmission and intercontinental spread. Science 327: 469–474.
48. Fierer N, Hamady M, Lauber CL, Knight R (2008) The influence of sex, handedness, and washing on the diversity of hand surface bacteria. Proc Natl Acad Sci U S A 105: 17994–17999.
49. Harris JK, De Groote MA, Sagel SD, Zemanick ET, Kapsner R, et al. (2007) Molecular identification of bacteria in bronchoalveolar lavage fluid from children with cystic fibrosis. Proc Natl Acad Sci U S A 104: 20529–20533.
50. Chao A, Lee S-M (1992) Estimating the number of classes via sample coverage. J Amer Stat Assoc 87: 210–217.
51. Chao A (1984) Nonparametric estimation of the number of classes in a population. Scand J Statist 11: 265–270.
52. R Development Core Team (2005) R: A Language and Environment for Statistical Computing. Vienna, Austria: R Foundation for Statistical Computing.
53. Oksanen J, Kindt R, Legendre P, O'Hara B, Simpson GL, et al. (2008) Vegan: Community Ecology Package. R package version 1.15-1. Available: http://cran.r-project.org/.
54. Kaufman L, Rousseeuw PJ (2005) Finding Groups in Data: An Introduction to Cluster Analysis. New York: Wiley.
55. Warner GR Gplots: Various R programming tools for plotting data. 2.6.0 ed. Available: http://cran.r-project.org/web/packages/gplots/index.html.