Molecular analysis of bacterial contamination on stethoscopes in an intensive care unit

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Abstract

Background: Culture-based studies, which focus on individual organisms, have implicated stethoscopes as potential vectors of nosocomial bacterial transmission. However, the full bacterial communities that contaminate in-use stethoscopes have not been investigated.

Methods: We used bacterial 16S rRNA gene deep-sequencing, analysis, and quantification to profile entire bacterial populations on stethoscopes in use in an intensive care unit (ICU), including practitioner stethoscopes, individual-use patient-room stethoscopes, and clean unused individual-use stethoscopes. Two additional sets of practitioner stethoscopes were sampled before and after cleaning using standardized or practitioner-preferred methods.

Results: Bacterial contamination levels were highest on practitioner stethoscopes, followed by patient-room stethoscopes, whereas clean stethoscopes were indistinguishable from background controls. Bacterial communities on stethoscopes were complex, and community analysis by weighted UniFrac showed that physician and patient-room stethoscopes were indistinguishable and significantly different from clean stethoscopes and background controls. Genera relevant to healthcare-associated infections (HAIs) were common on practitioner stethoscopes, among which Staphylococcus was ubiquitous and had the highest relative abundance (6.8%–14% of contaminating bacterial sequences). Other HAI-related genera were also widespread although lower in abundance. Cleaning of practitioner stethoscopes resulted in a significant reduction in bacterial contamination levels, but these levels reached those of clean stethoscopes in only a few cases with either standardized or practitioner-preferred methods, and bacterial community composition did not significantly change.

Conclusions: Stethoscopes used in an ICU carry bacterial DNA reflecting complex microbial communities that include nosocomially important taxa. Commonly used cleaning practices reduce contamination but are only partially successful at modifying or eliminating these communities.

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Stethoscopes are frequently used on multiple patients, and they have been implicated as vectors for nosocomial transfer of bacteria responsible for healthcare-associated infections (HAIs). It is well documented that practitioner stethoscopes are not routinely disinfected,1,2 and studies based on bacterial culture show that they may be contaminated with potential pathogens including methicillin-resistant and -sensitive Staphylococcus spp, multidrug-resistant P. aeruginosa, Acinetobacter spp, Enterococcus spp, Escherichia coli, Klebsiella spp, and Streptococcus spp.3–5 Culture-based studies have also shown that thorough stethoscope decontamination can significantly reduce pathogen colony-forming units (CFUs),6,7 although the impact of actual practitioner practices is less clear. Culture-based studies are limited, however, because culture can only identify agents of a priori interest but not entire microbial communities that may be present.

In contrast to culture, which is focused on individual bacteria and is only semi-quantitative, emerging molecular approaches using next-generation sequencing can provide unbiased profiling of entire bacterial communities in a manner that is both comprehensive and highly quantitative.8,9 These powerful approaches have revolutionized studies of the microbiome and of microbial ecology. Here, we used next-generation sequencing to investigate bacterial contamination on several types of stethoscopes in a medical intensive care unit (ICU), including stethoscopes carried by practitioners and used with multiple patients. We also investigated the effects of cleaning protocols that are used by practitioners in everyday practice.

Materials and Methods

Sample collection method (set A)

Stethoscope samples were collected in the medical ICU at the Hospital of the University of Pennsylvania. Stethoscope diaphragms were swabbed for 60 seconds using a flocked swab.
moistened with sterile saline (Copan Diagnostics, Murrieta, CA). Set A included swabs collected from 10 single-use disposable stethoscopes directly from the box prior to use (clean stethoscopes), 20 single-use disposable stethoscopes in-use in inpatient rooms (patient-room stethoscopes), and 20 stethoscopes being carried by physicians, nurses, and respiratory therapists (practitioner stethoscopes). Because low levels of microbial DNA are ubiquitous and bacterial DNA derived from collection instruments, reagents or the environment can confound microbiome studies,10,11 a set of background controls comprised of swabs moistened with saline (collected in parallel with the stethoscope sampling) were obtained on each collection date (n = 20). Swabs were stored at −80°C.

**Standardized cleaning method (set B)**

We sampled 10 additional ICU practitioner stethoscopes using the procedure described above, following which stethoscope diaphragms were cleaned with a hydrogen peroxide wipe (hydrogen peroxide 1.4%; Clorox Healthcare, Oakland, CA) for 60 seconds and allowed to dry. Stethoscopes were then swabbed again using the same protocol.

**Practitioner preferred cleaning method (set C)**

An additional 20 practitioner stethoscopes were collected. To ensure that the pre-cleaning swab would not affect the post-cleaning communities, the stethoscope diaphragm was sampled as described above, but only the left half of the diaphragm was swabbed. Stethoscopes were returned to the practitioner, and they were instructed to clean their stethoscopes using the method they usually would use to clean it between patients. Practitioners cleaned their stethoscopes with hydrogen peroxide wipes (n = 14), alcohol swabs (70% isopropyl alcohol; Coviden-Webcol, Mansfield, MA) (n = 3), or bleach wipes (sodium hypochlorite 0.55%; Clorox Healthcare) (n = 3), and duration of cleaning was determined by practitioner preference. Once the diaphragm was visibly dry, the right half of the diaphragm was then swabbed to capture post-cleaning bacterial communities.

**Bacterial extraction**

Swabs were cut directly into PowerSoil beadbeater tubes (MoBio, Qiagen, Venlo, Netherlands). DNA was extracted using the PowerSoil DNA kit (MoBio, according to the manufacturer’s instructions except for an additional 10-minute, 95°C incubation step to improve DNA yield from hard-to-lyse bacteria. DNA was stored at −20°C.

**Bacterial amplification sequencing and analysis**

Extracted DNA was amplified in triplicate 25-μL reactions using barcode-labeled primers targeting the bacterial 16S rRNA gene variable regions 1 and 2 (V1V2), employing previously described polymerase chain reaction (PCR) primers and protocols.12,13 The PCR primers target sequences that are conserved among bacteria, whereas the region amplified is not conserved and provides sequence-based information for bacterial identification. Samples were purified using bead purification, quantified using PicoGreen (Fisher Scientific, Waltham, MA), normalized to 5 ng/μL per sample, pooled, and subjected to deep sequencing on the Illumina MiSeq platform. Clean swabs moistened in sterile saline and dry clean swabs were run in parallel as controls for instrument- or reagent-derived background sequences. Using the QIIME 1.91 pipeline,14 sequences were clustered based on 97% similarity into de novo operational taxonomic units (OTUs), which serve as the basic taxonomic unit for subsequent analysis. To identify the bacteria, OTUs were aligned to the Greengenes reference database of bacterial sequences. Stethoscope set C was also amplified using primers targeting the 16S rRNA gene V4 region15 and was sequenced and analyzed using the same protocol. Sequences aligning with Streptophyta, which represent chloroplast DNA, were removed from the analysis.16

Bacterial DNA was quantified using 2 methods. First, the amount of amplification product generated by bar-coded PCR primer amplification during library preparation was used to estimate the relative amount of contamination in pre- and post-cleaning specimens, as previously described.17 In addition, 16S rRNA gene qPCR was carried out on a subset of samples using primers and protocols previously described.18

To identify bacteria (taxa) that are commonly associated with HAIs, we specifically queried sequences from practitioner stethoscope samples of sets A and C for the presence of *Staphylococcus*, *Pseudomonas*, *Acinetobacter*, *Closstridium*, *Enterococcus*, *Stenotrophomonas*, and *Burkholderia* genera. The 16S rRNA gene sequences assigned to these genera in the QIIME pipeline were then manually aligned to the NCBI 16S rRNA sequence database with BLAST to confirm genus identity and, if possible, to generate species-level assignment. Because any 16S rRNA gene primer set may have unrecognized biases,19,20 we did this using both V1V2 and V4 sequences of the 16S rRNA gene. Any sample with ≥10 sequence reads aligning to the genus was considered a positive hit.

**Statistical analysis**

Figures were generated and statistical tests carried out using R version 3.2.3 software (R Foundation for Statistical computing, Vienna, Austria). Richness (number of taxa) and alpha diversity (within-community measure that encompasses both richness and evenness) was carried out after sequence rarefaction to 1,000 reads.21 Alpha diversity was calculated using the vegan package. The pairwise Wilcoxon rank-sum test was used to compare between-group differences for the alpha diversity analysis. We analyzed β diversity (a metric of between-community differences) was analyzed by unweighted and weighted UniFrac using the QIIME pipeline, and was used to perform principal coordinate analysis (PCoA).14,22 The adonis function in the vegan package was used to test for statistical significance between groups of communities using PERMANOVA in the β diversity analyses. The Wilcoxon rank-sum test was used to examine the differences between groups of communities in quantification, richness and diversity, and the Student t test was used for the pre- and post-cleaning 16S quantification differences.

**Results**

**Stethoscope microbiome community analysis**

We first estimated total bacterial contamination of practitioner stethoscopes, patient-room stethoscopes, clean unused stethoscopes, and background controls in set A according to the quantity of 16S amplicon post-PCR amplification (Fig. 1(A)). Practitioner stethoscopes had significantly higher 16S amplicon concentration compared to both the patient-room and clean stethoscopes (P = .035 and P = .004, respectively; Wilcoxon rank-sum test). Patient-room stethoscopes had significantly higher
concentrations than the clean stethoscopes ($P = 1.8 \times 10^{-5}$; Wilcoxon rank-sum test). Both practitioner and patient-room stethoscopes were significantly higher in 16S quantity than background controls, whereas clean stethoscopes were indistinguishable from the background controls ($P = .967$; Wilcoxon rank-sum test).

We analyzed species richness, which reflects the number of different taxa within each community (Fig. 1(B)). Practitioner stethoscopes had significantly greater richness than either patient-room or clean stethoscopes ($P = .003$ and $P = .004$, respectively; pairwise Wilcoxon rank-sum test). All stethoscope groups were significantly higher in richness than the background controls. There was no significant difference in richness between patient-room and clean stethoscopes.

We then assessed $\alpha$ diversity of the stethoscope bacterial communities using the Shannon diversity index, a metric that incorporates both richness and evenness of distribution, with higher diversity reflecting greater richness and more even distribution of taxa (Fig. 1(C)). Practitioner, patient-room, and clean stethoscopes were all significantly more diverse than the background controls ($P = .0005$, $P = .003$, and $P = .0004$, respectively; pairwise Wilcoxon rank-sum test). In contrast, there was no significant difference in Shannon diversity between stethoscope groups. Taxa identified on practitioner stethoscopes at $>1\%$ relative abundance within their respective communities are shown in a heatmap in Fig. S1.

To compare the overall bacterial communities of the different groups, we calculated UniFrac distances among samples using unweighted and weighted methods, then plotted them on principal coordinates analysis (PCoA) plots (Fig. 2). The UniFrac metric compares complex microbial communities based on the phylogenetic relatedness of the bacteria contained within the communities, and the PCoA plot provides an overview visualization of community relatedness.

Using the unweighted UniFrac metric (Fig. 2(A)), which takes into account the presence or absence of taxa in different samples but not their relative abundance, bacterial communities were significantly different between all stethoscope groups ($P < .001$; PERMANOVA). Using the weighted UniFrac metric (Fig. 2(B)), which takes into account the presence or absence as well as relative abundance of bacteria comprising the communities, we observed that practitioner and patient-room stethoscopes differed from the clean stethoscopes and background controls ($P < .001$; PERMANOVA), but the practitioner and patient-room stethoscopes were not significantly different from one another ($P = .106$; PERMANOVA), nor were clean stethoscope and background controls ($P = .07$; PERMANOVA). These results suggest that both types of in-use stethoscopes differ substantially from the 2 types of control samples (clean stethoscopes and background), and that low-abundance taxa were mainly responsible for the differences between practitioner and patient-room stethoscopes, and for the differences between clean stethoscopes and background controls.

We next sought to determine which genera were responsible for the differences between bacterial communities on the in-use stethoscopes and the 2 types of controls on the weighted UniFrac PCoA. Figure 3 shows the top taxa responsible for separation of the communities plotted as vectors on the weighted PCoA, thus indicating which bacteria are responsible for differentiating communities located in distinct regions of the PCoA. The genera *Methylobacterium*, *Pseudomonas*, and *Acinetobacter* drove the separation of the clean stethoscopes and background controls from the practitioner and patient-room stethoscopes, indicating that these taxa are mainly derived from background sources in this sample set. Conversely, the practitioner and patient-room samples were characterized by *Porphyromonas*, *Bacteroides*, *Granulicatella*, *Actinomyces*, *Prevotella*, *Streptococcus*, *Staphylococcus*, *Corynebacterium*, and *Propionibacterium*, which are common oral, skin, and gut bacteria.

Effects of cleaning on stethoscope bacterial biomass and communities

Although patient-room stethoscopes are typically used only for a single patient, practitioner stethoscopes are used on multiple patients, raising the possibility of microbial transfer. Because practitioners may clean their stethoscopes between uses, we analyzed the effect of cleaning on the bacterial biomass on stethoscope diaphragms based on levels of 16S rRNA DNA. For 1 set of stethoscopes (set B; $n = 10$), we used a standardized cleaning method: vigorously wiping the diaphragm with a hydrogen-peroxide wipe for 60 seconds. For another set (set C; $n = 20$), we asked each practitioner to clean the diaphragm themselves with the method they
usually use between patients. Bacterial contamination was quantified by the amount of 16S rRNA gene amplicon following barcoded PCR quantification was done by 16S qPCR (Fig. S2).

We then compared the bacterial community composition of practitioner Set C before and after practitioner cleaning (Fig. 5). In the unweighted Unifrac PCoA, which is equally impacted by high- and low-abundance taxa (Fig. 5(A)), the pre- and post-cleaning samples were significantly different from one another ($P = .001$; PERMANOVA) and were significantly different from the clean stethoscopes and background controls ($P < .001$; PERMANOVA). In contrast, no significant difference was observed between the pre- and post-practitioner cleaning in the weighted Unifrac PCoA, in which taxa are weighted based on their relative abundances ($P = .274$; PERMANOVA) (Fig. 5(B)). These results suggest that cleaning does not have a substantial effect on community structure and that low abundance (minority) taxa are mainly responsible for differences between the pre- and post-cleaning communities seen in the unweighted but not weighted analysis.

Identification of nosocomial genera

We next investigated the presence of bacteria that are potential nosocomial pathogens. Samples from practitioner stethoscopes in sets A and C (prior to cleaning) were queried for relevant bacteria (Table 1). Both the V1V2 and V4 16S rRNA gene-variable regions were targeted to minimize the impact of potential primer biases.$^{19,20}$ In addition to aligning 16S rRNA gene sequences through the QIIME pipeline, sequences were manually aligned to databases by BLAST to optimize assignment.

Sequences were identified from genera that are commonly associated with hospital-associated infections including *Staphylococcus*, *Pseudomonas*, *Acinetobacter*, *Clostridium*, *Enterococcus*, *Stenotrophomonas*, and *Burkholderia*. Most taxa could only be assigned at the genus level by both V1V2 and V4 16S rRNA gene sequences, although some *Staphylococcus* sequences could be identified at the species level and included *S. aureus*. All stethoscopes had *Staphylococcus* spp., and more than half of them were confirmed to carry *S. aureus*. Most stethoscopes also carried *Pseudomonas* and *Acinetobacter*. Approximately half of the stethoscopes had *Enterococcus*, *Stenotrophomonas*, and *Clostridium*, while *Burkholderia* was less frequent. The V4 results were overall highly concordant with
V1V2 sequences, although the V4 region detected a somewhat higher prevalence of *Burkholderia* and *Clostridium* and somewhat less *Stenotrophomonas*. For each of these genera, we calculated the relative abundance (i.e., proportion of all bacterial sequences in a sample that were assigned to that genus) across all stethoscopes. *Staphylococcus* was present at 6.8%–14% relative abundance in the 2 practitioner stethoscope sets based on V1V2 and V4 sequences. *Pseudomonas* and *Acinetobacter* were less abundant and were present at slightly greater than 1% based on V4 sequences and ~10-fold lower based on V1V2 sequences, whereas other genera were below 1% in abundance. Thus, practitioner stethoscopes that are used on multiple patients carry sequences of genera that include important nosocomial pathogens.

**Discussion**

This study is the first to apply comprehensive molecular profiling to understand stethoscope contamination in a healthcare setting. We found that stethoscopes carried by practitioners in an ICU, which are used on multiple patients, are significantly contaminated with a rich and diverse community of bacteria that includes genera associated with HAIs. We also examined the impact of cleaning and found a surprisingly modest effect on contaminating bacterial communities. Taxa of skin, gut and oral sources dominated stethoscope bacterial communities, and genera associated with nosocomial infection were common. *Staphylococcus* was not only ubiquitous (found on 40 of 40 stethoscopes tested) but was also present at high abundance, representing 6.8%–14% of all bacterial sequences, depending on stethoscope set and target region queried. Most OTUs could not be assigned at the species level with either V1V2 or V4 sequences. However, definitive *S. aureus* assignment was possible for sequences on 24 of 40 practitioner stethoscopes. Therefore, at a minimum, more than half of these stethoscopes were contaminated by *S. aureus*. *Pseudomonas* and *Acinetobacter* were also widely present, with the exact frequency and relative abundance differing depending on the 16S variable region sequenced. Different 16S primer sets can have biases in the efficiency of detecting specific sequences, and our findings using...
2 different sets clearly establish that these taxa are widely present on stethoscopes, albeit at low relative abundances. Other nosocomial infection-related genera were also identified on practitioner stethoscopes at high frequency but lower relative abundances. Because this is the first study to analyze stethoscope contamination at the molecular level, it remains to be determined what amount of contamination is clinically relevant for potential nosocomial transmission.

We also applied these molecular methods to determine the impact of cleaning the stethoscope diaphragm, testing both a standardized approach and practitioners’ usual methods. Both methods significantly reduced the 16S DNA bacterial biomass but failed to bring contamination to the level of clean stethoscopes. The standardized cleaning method reduced more stethoscopes to the clean level (5 of 10) than the practitioner-preferred method (2 of 20). PCoA analysis of bacterial communities before and after cleaning revealed a shift toward background communities detected by the analysis of community membership only, but not when bacterial relative abundance was incorporated (unweighted vs weighted Uni-Frac; Fig. 5). This results implicates low-abundance taxa as contributing the most to differences, whereas the more abundant taxa are not substantially altered by cleaning. Notably, although the CDC offers recommendations on stethoscope decontamination, 2 studies suggest that it is infrequently practiced.1,2

Our study has several limitations. We were not able to identify bacterial taxa at the species level for most OTUs in our data set, and thus most HAI-associated bacteria could only be defined at the genus level. Future studies might complement comprehensive unbiased 16S rRNA gene analysis with species-targeted amplification. We were not able to identify drug-resistant species using this method. Because this is a DNA-based approach, it cannot distinguish between bacteria that are dead versus alive.24 Finally, we did not analyze fungal or viral sequences to understand the complete scope of organisms that stethoscopes carry in the ICU.

In summary, this study is the first comprehensive molecular analysis of bacterial contamination of stethoscopes in an ICU, the presence of potential nosocomial pathogens, and the impact of cleaning methods. Practitioner stethoscopes are contaminated by a plethora of bacteria, including organisms that may be associated with nosocomial infections. Cleaning reduces contamination but does not bring the bacterial biomass down to the level of clean stethoscopes nor does it significantly change the overall community composition. Thus, stethoscopes are a potential vector of HAI transfer. Useful future directions would be to use these molecular approaches to identify improved cleaning methods, enhance species-level identification of pathogens, quantify live versus dead bacteria, and define fungal and viral contaminants as well. In addition, shotgun metagenomic sequencing would be useful to analyze drug-resistance genes that might be carried between patients on practitioner stethoscopes.

**Table 1.** Identification of Potential Nosocomial Pathogens on Practitioner Stethoscopes

<table>
<thead>
<tr>
<th>Genus</th>
<th>Set A (n = 20) V1V2 Region</th>
<th>Set C (n = 20) V1V2 Region</th>
<th>V4 Region</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Frequency, No. (%)</td>
<td>Relative Abundance (mean ± SD)</td>
<td>Frequency, No. (%)</td>
</tr>
<tr>
<td><em>Staphylococcus</em> (g)</td>
<td>20 (100)</td>
<td>6.82 ± 5.00</td>
<td>20 (100)</td>
</tr>
<tr>
<td><em>S. aureus</em> (s)</td>
<td>11 (55)</td>
<td></td>
<td>13 (65)</td>
</tr>
<tr>
<td><em>Pseudomonas</em> (g)</td>
<td>16 (80)</td>
<td>0.23 ± 0.27</td>
<td>20 (100)</td>
</tr>
<tr>
<td><em>Acinetobacter</em> (g)</td>
<td>13 (65)</td>
<td>0.14 ± 0.14</td>
<td>20 (100)</td>
</tr>
<tr>
<td><em>Clostridium</em> (g)</td>
<td>8 (40)</td>
<td>0.06 ± 0.13</td>
<td>12 (60)</td>
</tr>
<tr>
<td><em>Enterococcus</em> (g)</td>
<td>8 (40)</td>
<td>0.04 ± 0.06</td>
<td>18 (90)</td>
</tr>
<tr>
<td><em>Stenotrophomonas</em> (g)</td>
<td>7 (35)</td>
<td>0.03 ± 0.05</td>
<td>18 (90)</td>
</tr>
<tr>
<td><em>Burkholderia</em> (g)</td>
<td>3 (15)</td>
<td>0.01 ± 0.01</td>
<td>3 (15)</td>
</tr>
</tbody>
</table>

Note. SD, standard deviation.

Sequence data from the practitioner stethoscopes in sets A and C were investigated for the presence of selected bacterial genera (g) that commonly cause hospital-acquired infections (HAIs), based on 16S rRNA gene V1V2 or V4 region sequences as described in the Methods section. Frequency indicates the proportion of practitioner stethoscopes with that genus. Relative abundance indicates the percentage of all bacterial sequences on a stethoscope that genus represents, across the practitioner stethoscope sets. For *Staphylococcus*, some sequences could be assigned at the species level (s), and those identified as *S. aureus* are indicated.

**References**


**Supplementary materials.** To view supplementary material for this article, please visit https://doi.org/10.1017/ice.2018.319

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