Exploring Bacterial Diversity in Hospital Environments by GS-FLX Titanium Pyrosequencing

Margarita Poza1*, Carmen Gayoso1, Manuel J. Gómez2, Soraya Rumbo-Feal1, María Tomás1, Jesús Aranda1, Ana Fernández1, Germán Bou1*

1 Microbiology Department, Biomedical Research Institute-University Hospital, A Coruña, Spain, 2 Sequencing and Bioinformatics Department, Astrobiology Center INTECSIC, Madrid, Spain

Abstract

Understanding microbial populations in hospital environments is crucial for improving human health. Hospital-acquired infections are an increasing problem in intensive care units (ICU). In this work we present an exploration of bacterial diversity at inanimate surfaces of the ICU wards of the University Hospital A Coruña (Spain), as an example of confined hospital environment subjected to selective pressure, taking the entrance hall of the hospital, an open and crowded environment, as reference. Surface swab samples were collected from both locations and recovered DNA used as template to amplify a hypervariable region of the bacterial 16S rRNA gene. Sequencing of the amplicons was performed at the Roche 454 Sequencing Center using GS-FLX Titanium procedures. Reads were pre-processed and clustered into OTUs (operational taxonomic units), which were further classified. A total of 16 canonical bacterial phyla were detected in both locations. Members of the phyla Firmicutes (mainly Staphylococcus and Streptococcus) and Actinobacteria (mainly Micrococcaceae, Corynebacteriaceae and Brevibacteriaceae) were over-represented in the ICU with respect to the Hall. The phylum Proteobacteria was also well represented in the ICU, mainly by members of the families Enterobacteriaceae, Methylobacteriaceae and Sphingomonadaceae. In the Hall sample, the phyla Proteobacteria, Bacteroidetes, Deinococcus-Thermus and Cyanobacteria were over-represented with respect to the ICU. Over-representation of Proteobacteria was mainly due to the high abundance of Enterobacteriaceae members. The presented results demonstrate that bacterial diversity differs at the ICU and entrance hall locations. Reduced diversity detected at ICU, relative to the entrance hall, can be explained by its confined character and by the existence of antimicrobial selective pressure. This is the first study using deep sequencing techniques made in hospital wards showing substantial hospital microbial diversity.

Citation: Poza M, Gayoso C, Gómez MJ, Rumbo-Feal S, Tomás M, et al. (2012) Exploring Bacterial Diversity in Hospital Environments by GS-FLX Titanium Pyrosequencing. PLoS ONE 7(8): e44105. doi:10.1371/journal.pone.0044105

Editor: Matthias Horn, University of Vienna, Austria

Received February 1, 2012; Accepted July 30, 2012; Published August 29, 2012

Copyright: © 2012 Poza et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This work was supported by the Spanish Network for Research in Infectious Diseases (REIPI, RD06/0008/0025), grants from Fondo de Investigaciones Sanitarias (PI081368 and PS09/00687), SERGAS (PS07/90) and Xunta de Galicia (07CSA050916PR) to GB and a grant from SERGAS (PS08/24, Galicia, Spain) to MP. MP was supported by the Isidro Parga Pondal Program (Xunta de Galicia, Spain), CG by the Spanish Network for Research in Infectious Diseases (REIPI, RD025/0008), MUG by the project Consolider INGENIO CS02007-0005 and INTA-CSIC (Madrid, Spain) and SRF, MT, JA and AF by Instituto de Salud Carlos III (Spain). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: German.Bou.Arevalo@sergas.es

These authors contributed equally to this work.

Introduction

Metagenomics advances due to the increased availability of high throughput platforms for DNA sequencing and associated bioinformatics software are revolutionizing our knowledge about microbial communities. Direct studies of microorganisms that are difficult to culture (as much as 99% of the microorganisms living in a natural environment) may now be performed by these modern techniques. Prokaryotic 16S ribosomal RNA genes (16S rRNA genes in the case of eukaryota) have been commonly used for studying biodiversity since they are universally distributed in all cellular organisms and allow the identification and comparison of microorganisms present in a wide variety of environmental samples [1–3]. Many authors have used this approach to explore 16S rRNA diversity in natural samples [4–6]. Pyrosequencing techniques have the great advantage of allowing direct sequencing of 16S rRNA collections without the need of the cloning step. The obtained collection of 16S rRNA sequences can then be compared against reference databases such as GenBank and RDP (Ribosomal Database Project, http://rdp.cme.msu.edu/), which include more than 800,000 16S rRNA sequences, with the aim of having them classified.

In the last decade, many studies have been performed to describe microbial populations of samples ranging from the abyssal sea floor [7] or the polar desert [8] to the water and air of a hospital therapy pool [9]. In 2004, whole genome shotgun sequencing techniques were applied to microbial populations collected from sea water samples of the Sargasso Sea [10]. Since then, many studies describing microbial diversity have been done applying deep sequencing technologies from different samples as marine environments [11,12] or forest and grassland soils [6]. In only one decade it became clear that most of the bacteria isolated from natural environments were not represented in culture collections. Moreover, recent studies directed to describe microbial populations living in the human body, as the human skin microbiome characterized by Grice et al., [4] or the oral or gastrointestinal...
microbiome, explored by several groups using deep sequencing technologies [13–19], have revolutionized our understanding about the substantial diversity of microorganisms co-habiting with us. The application of new technologies has promoted the discovery that biodiversity is much more complex than traditional culture methods indicated in the past.

Hospital-acquired infections are the sixth leading cause of death in the United States and similar data have been reported from Europe [20], being an increasing problem in intensive care units (ICU), where the patients are more susceptible to colonization and the organisms are often more resistant to antimicrobial agents than in other environments. Since the 80’s, infectious disease specialists have recognized that ICU patients acquire nosocomial infections at a much higher rate than patients elsewhere in the hospital [21]. In these confined areas, lower respiratory tract and bloodstream infections happen to be the most lethal; however, urinary tract infections are the most common. Infections caused by gram-negative bacteria have features that are of particular concern since these organisms are highly efficient at up-regulating or acquiring genes that code for mechanisms of antibiotic drug resistance, especially in the presence of antibiotic selection pressure. Most common nosocomial pathogens may well survive or persist on surfaces for a long time, what could explain their survival in ICU wards [22]. Gram-positive bacteria, such as Enterococcus spp. (including VRE), Staphylococcus aureus (including MRSA), Clostridium difficile or Streptococcus pyogenes, and many gram-negative bacteria, such as Acinetobacter spp., Klebsiella spp., Pseudomonas aeruginosa, Seratia marcescens, or Shigella spp., may indeed survive for months on dry surfaces. A few others, such as Bordetella pertussis, Haemophilus influenzae, Proteus vulgaris, or Vibrio cholerae, however, persist only for days. In the present work we describe for the first time the substantial diversity of microorganisms co-habiting with us. The application of new technologies has promoted the discovery that biodiversity is much more complex than traditional culture methods indicated in the past.

Results

Sequence Data

A fragment of the V7–V9 hypervariable region of the bacterial 16S rRNA gene, around 450–480 base pairs long, was amplified by PCR with primers mp1682f and mp1682r from genomic DNA obtained from surface swab samples collected at ICU ward and at the entrance hall of the University Hospital A Coruña (Spain). The two collections of amplicons were subjected to 454 pyrosequencing at the Roche Sequencing Center. The number of reads obtained was 501707 and 346427 for ICU and Hall samples, respectively (Table 1).

Table 1. Total number of reads accumulated for each sample in the original data sets and after filtering and classification.

| Sample | Original number of reads | Number of reads after pre-processing | Number of reads after sub-sampling | Number of reads classified by Mothur |
|--------|--------------------------|-------------------------------------|-----------------------------------|-------------------------------------|
| ICU    | 501707                   | 95104 (18%)                         | 95104 (18%)                       | 95103 (18%)                         |
| Hall   | 346427                   | 99952 (28%)                         | 95104 (27%)                       | 95071 (27%)                         |

Sequence Pre-processing

The analysis of the read length distributions indicated that the modal length was 458 nt for the ICU sample and 457 for the Hall one. However, a significant number of reads had shorter lengths. Reads were filtered and trimmed as described in Materials and Methods, resulting in datasets containing around 42% and 43% of the original reads for the ICU and Hall samples, respectively. Their alignment, against the Core Set reference alignment of the Greengenes database, and subsequent screening, to optimize the number of them covering the same sequence space, resulted in datasets containing around 19% and 30% of the original reads. Reads were then grouped using a pseudo-single linkage algorithm, to reduce the impact of pyrosequencing noise, and screened with UCHIME to discover and eliminate chimeric sequences. This last step eliminated 7635 chimeric sequences from both samples, resulting in datasets containing 18% and 28% of the original reads for the ICU and Hall samples, respectively (Table 1). Finally, datasets were sub-sampled to normalize the number of reads in each sample (Table 1).

OTU Detection

Operational taxonomic units (OTUs) were defined by clustering pre-processed, grouped and normalized reads as described in Materials and Methods. The number of OTUs detected at a distance of 0.03, which is assumed to correspond to the species level, was 1145 for the ICU sample and 2499 for Hall sample (Table 2). The total number of OTUs observed in the two samples was 3000. The number of OTUs that were observed as shared between the ICU and Hall samples was 644 (Fig. 1). The Ace and Chao1 estimates of community richness at a distance of 0.03 had values of 2798 and 2117, respectively, for the ICU sample, and 5043 and 4279 for the Hall sample (Table 2), suggesting that diversity at species level could still be up to two times higher than observed, approximately, and that the Hall sample contained a higher diversity than the ICU sample, in terms of species richness. In agreement with this observation, the shape of rarefaction curves calculated at 0.03, 0.05 and 0.1 distances indicate a stronger trend to saturation for the ICU sample than for the Hall sample (Fig. 2). It should be noted that, to prevent the obliteration of rare phylotypes, singleton OTU removal was not included in the data processing pipeline. Therefore, observed and estimated richness values could be overestimated. However, this possibility would not be an explanation for the difference in richness observed for the Hall and ICU samples, since a test in which singleton OTUs were discarded yielded the same result: Hall was twice as diverse as ICU, in terms of species richness (data not shown).

OTU Classification

Representative sequences from the 3000 OTUs detected at a distance of 0.03, corresponding to a total of 190174 sequence reads from the two samples, were classified up to genus level, as described in Materials and Methods (see Dataset A in http://
www.cab.inta-csic.es/usb/HMG/. Supporting Information are interactive files therefore they have been included in an author’s website). Only 11 OTU representative sequences, corresponding to 34 individual reads, could not be classified (Table 1). Figure 3 presents a comparison of OTU relative abundances at the order level in both samples and Figure 4 shows the relative abundance of families detected in ICU and Hall samples. A hierarchical pie chart, representing the bacterial diversity recovered from ICU and Hall samples is presented in Figure 5, where the most representative genera are explicitly indicated. For more details, interactive representations can be found in Figure A, Figure B and Figure C (see http://www.cab.inta-csic.es/usb/HMG/).

To identify taxonomic nodes with differential presence in any of the two environments, observed frequencies were further normalized and relative differences (rd) were calculated for each taxonomic node. The statistical significance of the differences was independently assessed by computing a Chi-Square test. A total of 16 canonical phyla were detected in the present study, including both locations, and, as illustrated in Figure 5, two of them were over-represented in ICU with respect to the Hall: Firmicutes and Actinobacteria. The over representation of the phylum Firmicutes was due to the abundance of members of three orders: (i) Bacillales (rd = 63; p-value = 0), which was mainly consequence of the high abundance of members of the *Staphylococcus* genus (rd = 70; p-value = 0); (ii) Lactobacillales, mainly represented by *Streptococcus* (rd = 48; p-value = 0); and (iii) Clostridiales, mainly represented by the family Veillonellaceae (rd = 80; p-value = 3.2E-202). The over representation of the phylum Actinobacteria was consequence of the high number of members of the order Actinomycetales (rd = 51; p-value = 0) which was due to the abundant presence of members of the family Micrococcaceae (rd = 63; p-value = 0), as well as the family Corynebacteriaceae (rd = 55; p-value = 4.6E-289) and Brevibacteriaceae (rd = 96; p-value = 0). Other abundant group in ICU, although not over represented with respect to Hall, was phylum Proteobacteria. Proteobacteria were mainly represented in ICU by the family Enterobacteriaceae and by four orders that were indeed over represented in ICU with respect to Hall; (i) Rhizobiales (rd = 79; p-value = 0), specially members of the *Methylbacterium* genus (rd = 93; p-value = 0); (ii) Sphingomonadales, mainly from the *Sphingomonas* genus (rd = 78; p-value = 0); (iii) Pasteurellales (rd = 63; p-value = 1.33E-74); and (iii) Vibrionales (rd = 27; p-value = 3.1E-25). The phyla Proteobacteria, Bacteroidetes, Deinococcus-Thermus and Cyanobacteria were over represented in the Hall relative to ICU. The phylum Proteobacteria (rd = −36; p-value = 0) was highly represented in the Hall as consequence of the abundance of Enterobacteriales (rd = −54; p-value = 0), mainly as result of the predominant presence of unclassified members of the Enterobacteriaceae family (rd = −54; p-value = 0). Other orders of Proteobacteria over represented in the Hall were: (i) Burkholderiales (rd = −81; p-value = 0), specially the families Oxalobacteraceae (rd = −90; p-value = 0) and Comamonadaceae (rd = −42.73E-15; p-value = 0); (ii) Rhodobacteriales (rd = −45; p-value = 2.79E-72), mainly represented by the *Paracoccus* genus and (iii) Rhodospirillales (rd = −72; p-value = 2.63E-113). The phylum Bacteroidetes was
over represented in the Hall mainly because of the abundance of Sphingobacterales \((rd = -83; p-value = 0)\), especially members of the *Hymenobacter* genus \((rd = -83; p-value = 1.55E-299)\) and, in a lesser degree, to the order Flavobacterales \((rd = -64; p-value = 5.09E-238)\). The phylum Deinococcus-Thermus was basically represented by the order Deinococcales, mainly from the *Deinococcus* genus \((rd = -88; p-value = 9.68E-167)\). Finally, the over representation of the phylum Cyanobacteria \((rd = 87; p-value = 1.81E-297)\) was mainly due to unclassified bacteria. For more details see Table A (http://www.cab.inta-csic.es/usb/HMG/). Although not over represented with respect to ICU, Firmicutes and Actinobacteria were also significantly present in the Hall. Among Firmicutes, the most abundant orders were Bacillales, mainly represented by the *Staphylococcus* genus, and

| Sample | Reads | Clustering distance | OTU | ACE | Chao1 | OTU | ACE | Chao1 |
|--------|-------|---------------------|-----|-----|-------|-----|-----|-------|
| ICU    | 95104 | 0.03                | 1145| 2798| 2117  | 744 | 1729| 1343  |
| Hall   | 95104 | 0.05                | 2499| 5043| 4279  | 1636| 3206| 2696  |

Observed richness is presented, for each location, as the number of OTUs defined at clustering distances of 0.03, 0.05 and 0.1. Predicted richness is presented as the values of the Ace and Chao1 diversity estimators, at the same clustering distances.

**Table 2.** Observed and predicted richness.

**Figure 3.** Normalized abundance of reads associated with taxonomic nodes at the order level, expressed as the number of occurrences in 100000 reads. Abundances are shown in logarithmic (left panel) and decimal (right panel) scales.

doi:10.1371/journal.pone.0044105.g003
Lactobacillales, mainly *Streptococcus*. Among Actinobacteria, the most abundant order was Actinomycetales, mainly represented in the Hall by the genera *Corynebacterium* and *Propionibacterium* (for more information see Dataset A in http://www.cab.inta-csic.es/usb/HMG/). In general, the Hall sample was associated to a higher diversity than that from ICU, in agreement with richness estimations (Ace and Chao1 indices), which were approximately two times higher for the first location (Table 2). However, while the Hall was dominated by Enterobacteriaceae, plus a high number of taxa represented in moderate percentages, the ICU sample included less taxa that were more abundantly represented (i.e. Corynebacteriaceae, Micrococcaceae, Sphingomonadaceae, Neisseriaceae or Pasteurellaceae), as shown in Dataset A (http://www.cab.inta-csic.es/usb/HMG/).

**Clinical Data from Patients in ICU Wards**

To recover information about infections caused in the ICU wards during the experimental period, data obtained from clinical isolates collected from ICU-hospitalized patients was analyzed. Most of the microorganisms isolated from clinical samples belonged to the phylum Proteobacteria, mainly from the classes Gammaproteobacteria and Betaproteobacteria (Figure 6 and see also Figure D in http://www.cab.inta-csic.es/usb/HMG/). Among Gammaproteobacteria, the predominant groups were the family Enterobacteriaceae, mainly members of the genera *Escherichia*, *Enterobacter*, *Proteus*, *Citrobacter*, *Klebsiella*, *Serratia*, *Morganella* and *Salmonella*, and the orders *Pyramidales*, *Pseudomonadales*, *Aeromonadales*, and *Pasteuriidae*, *Moraxellales*, *Staphylococcales*, *Streptococcales*, *Sphingomonadales*, *Neisseriaceae* or *Pasteurellaceae*), as shown in Dataset A (http://www.cab.inta-csic.es/usb/HMG/).

**Discussion**

Microbiology has experienced a transformation during the last 30 years that has altered the microbiologists point of view about microbial populations. Metagenomics opened new fields of research by providing access to far more microbial diversity than has been ever viewed in the Petri dish [23]. Recently, advances in sequencing technologies have provided researchers the ability to sequence entire microbial communities in a high-throughput manner [24,25]. The relevance of the present study lies in the characteristics of the environments analyzed: inanimate surfaces at the ICU ward and the entrance hall of the University Hospital A Coruña. ICU wards are special and confined areas where selective pressure is extreme and cleaning measurements are strict, promoting the selection of microorganisms that can develop or acquire resistance mechanisms that allow them to survive in the presence of a broad spectrum of antimicrobial agents. Patients in ICU wards are critic, weak and immunosuppressed, in most cases, and therefore, even non virulent microorganisms may cause important infections leading to death. In contrast, the entrance hall of the hospital is an outward-facing and crowded place where a high diversity of people passes through and, therefore, where microbial diversity should be more related to that associated to the local human community. We consider that deep studies are needed to describe the microbial diversity associated to hospital environments, especially in ICU wards, to allow the identification of potential microorganisms causing nosocomial infections. A metagenomic study of the microbiota at hospital inanimate surfaces is difficult due to the small amount of DNA that can be recovered from the samples. This means that the genome sequences of abundant species will be well represented in a set of random shotgun reads, whereas lower abundance species may be represented by a small number of sequences. If sample size is reduced, less represented species could be missed. Pyrosequencing of 16S rRNA genes allows the deep characterization of complex microbial communities. Bacterial 16S rRNA gene contains nine hypervariable regions (V1–V9) that present considerable sequence diversity among different bacteria [26]. Combinations of conserved primer-binding sites and intervening variable sequences may facilitate genus and even species identification [27,28]. 454-Pyrosequencing techniques by GS-FLX Titanium chemistry protocols using primers amplifying the hypervariable domains V7–V9 were successfully used for the samples used in the present study. OTU detection showed that microbiota detected in ICU
was different from that in the Hall; both samples shared 644 OTUs of a total of 3000. We may extract from these data that microbial diversity differs in a confined ward subjected to selective pressure (ICU) with respect to an outward-facing and crowded hospital entrance hall, even taking into account that the lower diversity in ICU is due, in part, to the lower transit of people compared with the entrance hall, where thousands of individuals pass through a day. Besides, it is also important to remark that the area surface samples (around 200 m² ICU and 50 m² Hall) was not positively correlated with the diversity found. Although ICU ward is routinely strictly cleaned and supervised, a total of 15 canonical bacterial phyla were detected in this confined area in the present work. It was found that most of the microorganisms found in ICU belonged to the phyla Actinobacteria, Firmicutes, Proteobacteria and Bacteroidetes, that constituted 18.5%, 46.4%, 32.7%, and 1.4% of the bacterial community, respectively.
These data agree with previous results obtained for microbial communities found in the human body [18,29,30], what could be explained in terms of interactions between the normal microflora living on humans and the ICU ward environment. Important differences were addressed at the order, family and genera level between ICU and Hall samples, both in terms of abundance and diversity. Data showed that although many OTUs described were detected in both hospital samples, many others could only be detected in one of the environments, this justifying the view that the two microbiotas are different. Data from clinical samples collected from in-patients correlated well with pyrosequencing data since many genera over-represented in ICU were also represented in the clinical samples (i.e. *Staphylococcus* or *Streptococcus*, which are normal inhabitants of skin and mucosae, usually infects immunocompromised humans. As an extremely versatile pathogen, this genus has developed a broad spectrum of mechanisms leading to resistance to most powerful antimicrobial agents which could explain its presence in ICU wards. *S. aureus* is, in fact, one of the major causes of hospital acquired infection, being normal flora of ICUs [31]. The *Methylobacterium* members are methylotrophic bacteria that constitute part of the natural human foot and mouth flora. Certain species of *Methylobacterium* have been described in ICU wards causing bacteraemia in immunosupressed patients [32]. This genus showed much more abundance in ICU with respect to the Hall and its increased presence in ICU could be explained in terms of the methanol enriched agents routinely used for cleaning. *Sphingomonas* members are metabolically versatile and can use a wide range of naturally occurring compounds as carbon source, as well as some types of environmental contaminants. Its abundant presence in ICU could be explained because this genus has a good ability to utilize a wide range of organic compounds and to grow and survive under low-nutrient conditions, including toxic compounds [33]. As an example, *S. paucimobilis* has been previously detected in hospital equipment [34] and in respiratory therapy items, bedside water bottles, sinks…etc and has been described as a cause of minor infections in humans although it is also capable of causing serious and active infections [35]. *Streptococcus*, a diverse and versatile genus that causes a broad range of diseases, is highly over-represented in ICU with respect to Hall sample. In the last years, antibiotic-resistant *Streptococcus* strains have started appearing and causing epidemics [36,37]. *Corynebacterium* genus, normal flora of human skin, is found in a broad variety of habitats such as soil, plant material, food products and marine and animal sources. During the last years, a very considerable number of new corynebacterial species have been described as causing diseases to both healthy and immunocomprised hosts [38]. These data show that bacteria surviving in ICU must somehow be versatile...
microorganisms able to adapt in low nutritional conditions, to develop resistance mechanisms, to survive in the presence of toxic compounds or to persist in dry surfaces for a long time.

Strikingly, typical microorganisms naturally living in humid environments (i.e. *Cyanobacteria, Spirochaeta* or *Thiothrix*) have been detected in the Hall sample which could be due to the hospital location near the sea accompanied by the mild and damp climate of A Coruña.

All over the data showed that in the Hall sample there is much more biodiversity than in the ICU sample and that the most represented groups in ICU are more abundant in this ward than in the entrance Hall in many cases (i.e. *Streptococcus* or *Staphylococcus*). However, one of the most serious limitations of the present study lies on the sample size. It has to be taken into account that a bigger sample size including different hospital locations would improve or even modify the results. Even so, we could infer from the present study that the environment at ICU could be selecting for the presence of some species able to live in unfavourable conditions. We may conclude that the microbial biodiversity differs and it is reduced in a confined ward subjected to selective pressure (ICU) with respect to an outward-facing and crowded hospital entrance hall. Finally, in this study performed using new pyrosequencing technologies, we have found much more microbial diversity in hospital environments than ever seen before by classical methods. This study showed for the first time that hospital environments may act as filters selecting specific bacterial populations and creating special and confined areas where microbiota clearly differs from that present in nature.

Materials and Methods

Sample Collection

Samples were collected in three consecutive days of June 2009 at 8:00 a.m., before the routine cleaning, in the ICU ward (around 200 m²) and at the entrance hall (around 50 m²) of the University Hospital A Coruña, Spain. No specific permissions or ethics approval were required for these activities made at inanimate surfaces. No human samples were used. Sterile lints and gloves were used for swabbing all ICU surfaces. In the Boxes where patients are in bed, computer touch screens, monitors, drawers, infusion pumps, button panels of beds and respirator surfaces were swabbed. In the Pixies, computer touch screens and keyboards, doorknobs, drawers and gasometer surfaces were among the variety of sources from which samples were collected. In common areas and apparatus of the ICU wards, computer screens and keyboards, doors handles, cleaning room surfaces, medicine distributor surfaces, keyboard and electrodes of electrocardiographs, wrecker surfaces, knobs of refrigerators and drawers, microwaves and other surfaces were also cleaned with sterile lints.

In the case of the entrance hall, all the accessible surfaces were combed and cleaned using sterile lints.

Lints were introduced in sterile glass beakers (pre-treated with DEPC and sterilized 30 min, 121°C, 1.1 atm) containing sterile isotonic saline solution and incubated for 2–3 h at room temperature with gentle shaking. Then, lints were wringed out and the saline solution was filtered through 0.45 µm membranes (Millipore) and centrifuged 30 min at 4000 rpm in sterile tubes to recover the particulate fraction. At this point, all pellets from ICU and Hall samples were pooled into a single sample from each location.

Cell Wall Digestion and DNA Extraction

Both fraction pellets were resuspended in a buffer containing 20 mM Tris, pH 8, 2 mM EDTA and 1,2% Triton X-100. To digest the cell wall of the microorganisms supposedly present in the samples, the suspension was subjected to digestion using 10 mg/mL of lysozyme, 10 mg/mL of lysostaphin and 20 mg/mL of lyticase (all from Sigma-Aldrich). Digestions were performed for 1 h, at 37°C, in all cases. The Genomic Purification Kit (Promega) was then employed to recover total DNA from both samples, following the manufacturer’s instructions for each microbial type.

16S rRNA Amplification and Purification

In order to check the integrity of the 16S rRNA gene in the samples, oligonucleotides 16Sforw; 5’-AGA GTT TGA TCC TGG TCT AG-3’ and 16Srevr; 5’-GAC GGC GGG TGT GTR CA-3’, commonly used to amplify the complete 16S rRNA gene (ca. 1300 bp) as described by Grice et al., [4] were used for PCR. The reaction mixture contained 10 µL of template (DNA extracted as described above), 5 µL of a 20 mM solution of each oligonucleotide, 3 µL of 50 mM MgCl₂, 1 µL of a 40 mM of a mix of dNTPs, 5 µL of polymerase buffer, 0.5 µL of BioTaq polymerase (Bioline) and sterile water until 50 µL. The PCR program was as follows; one cycle of 95°C, 5 min followed by 20 cycles of 94°C, 30 s, 52°C, 30 s and 72°C, 1 min, and a final extension of 72°C, 2 min.

A segment of around 450–480 bp, containing the V7–V9 hypervariable regions of bacterial 16S rRNA gene [26] was amplified by PCR using oligonucleotides mfp16S2f/5’-GCA TGG ITG TCG TCA GCT CGT G and mfp16S2r/5’-AGC GIT ACC TTG TTA CGA GTT, using the primary DNA samples as template. The reaction mixture was as described above but using 1 µL of Expand High Fidelity polymerase (Roche). The PCR program was as follows; one cycle of 94°C, 2 min followed by 10 cycles of 94°C, 15 s, 52°C, 30 s and 72°C, 45 s, and 20 cycles of 94°C, 15 s, 52°C, 30 s and 72°C, 2 min 25 s, with a final extension of 72°C during 7 min. Controls were performed without template.

PCR products were purified using the MinElute Purification kit (Qiagen) being the DNA finally resuspended in ultrapure nuclelease-free water (Sigma-Aldrich).

The concentration and the purity grade of the samples were finally evaluated using a NanoDrop ND-1000 (Thermo scientific) and a BIOANALYZER 2100 employing the Agilent High Sensitivity DNA kit (both from Agilent Technologies, Inc. Germany).

Pyrosequencing

Sequencing of amplicons (ca. 2 µg of each pure sample) was conducted in the Roche 454 Sequencing Center (Connecticut, USA) using the GS-FLX Titanium chemistry and following the manufacturer’s protocol. Both ICU and Hall samples were sequenced in the same run.

Sequence Pre-processing and Sub-sampling

Most of the sequence processing and analyses procedures were carried out with Mothur [39], which is a package that combines a variety of tools designed to process 16S/18S rRNA gene sequence data and to characterize the diversity and structure of biological communities. Sequences were pre-processed by strictly following the Standard Operating Procedure (SOP) described by Schloss et al.[40]. In brief, pyrosequencing reads were first filtered and trimmed by imposing the following restrictions: maxambig = 0, maxhomop = 8, qwindowaverage = 35, qwindowsize = 50. This means that, those reads containing any ambiguity and those containing homopolymers longer than 8 nucleotides and, at the same time, trimmed the reads to the position in which the average quality score dropped below 33 over a 30-bp sliding window, were eliminated. Then, a non-redundant collection of sequence reads
was identified, to simplify the dataset, aligned against the Core Set reference alignment of the Greengenes database [41] and screened to ensure that most of the recovered reads aligned over the same sequence space. To reduce the impact of pyrosequencing noise, the resulting reads were grouped using the pseudo-single linkage algorithm pre.cluster, allowing a maximal number of 1% pair wise differences, following Mothur’s implementation of the pre-clustering method [42]. Sequences were then analyzed with UCHIME, to identify and remove chimeric reads, and classified to eliminate those that could be considered contaminants (mitochon-
drial, plastidial and archaeal sequences). Finally, sequences were sub-sampled to normalize the number of reads from each location.

**OTU Detection**

Pre-processed sequences, aligned and normalized, were used to generate an uncorrected pair wise distance matrix, which was then used to cluster the sequences using the furthest neighbor algorithm to detect OTUs at 0.03, 0.05 and 0.1 distances. Ace and Chaol estimates were calculated, to predict community richness, and rarefaction curves constructed, to assess sampling intensity.

**OTU Classification**

OTUs detected at a genetic distance of 0.03 were classified by comparing a representative sequence from each of them against the Silva reference database [43], using BLAST to identify similar sequences [44] and the k-Nearest Neighbor algorithm to determine the consensus taxonomy from the 10 most similar sequences in the database. Sequences were classified according to the RDP6 taxonomy scheme [45]. In general, this strategy allowed the classification of sequences into taxonomic categories that were never lower than genus.

**Identification of Differentially Distributed Taxonomic Nodes**

To identify taxonomic nodes for which a significant difference was observed in the number of reads associated to the ICU and Hall samples, the number of reads associated to each taxonomic node, for each of the samples, was further normalized by the total number of reads analysed classified by Mothur, after pre-processing and sub-sampling, for each sample. Then, for each taxonomic node, the relative difference (rd) between the numbers of reads classified in each sample was calculated and expressed as a percentage of the total number of reads classified in both samples for that taxonomic node. A Chi-Square test for homogeneity was also computed for each taxonomic node, to determine whether the normalized frequencies calculated for each sample were consistent with the null hypothesis that it is equally probable to detect a given taxon in any of the two samples.

**Clinical Data in ICU Wards**

A list of microorganisms isolated from clinical samples obtained from patients staying in ICU wards during the experimental period (May, June and July of 2009) was obtained from the records of the Microbiology Department of the University Hospital A Coruña. No specific permissions or ethics approval were required for these activities.

**Availability**

The pyrosequencing-derived 16S rRNA gene sequence datasets have been deposited in the GenBank short-read archive under accession number SRA046412.1.

**Acknowledgments**

Authors wish to express their gratitude to the ICU personnel from the University Hospital of A Coruña for their kindly help in sample collecting.

**Author Contributions**

Conceived and designed the experiments: MP CG MJG GB. Performed the experiments: MP CG SRF MT JA AF. Analyzed the data: MP CG MJG. Contributed reagents/materials/analysis tools: MJG. Wrote the paper: MP CG MJG GB.

**References**

1. Woese CR. (1987) Bacterial evolution. Microbiol Rev 51: 221–271.
2. Woese CR, Fox GE. (1977) Phylogenetic structure of the prokaryotic domain: the primary kingdoms. Proc Natl Acad Sci USA 74: 5089–5090.
3. Woese CR, Winker S, Gotell RR. (1990) Architecture of ribosomal RNA constraints on the sequence of “teta-loops”. Proc Natl Acad Sci USA 87: 5847–5847.
4. Grice EA, Kong HH, Coulan S, Deming CB, Davis J, et al. (2009) Topographical and temporal diversity of the human skin microbiome. Science 324: 1190–1192.
5. Armougom F, Raoult D. (2008) Use of pyrosequencing and DNA barcodes to monitor variations in Firmicutes and Bacteroidetes communities in the gut microbiota of obese humans. BMC Genomics 9: 576.
6. Nacke H, Thürmer A, Wollherr A, Will C, Hodac L, et al. (2011). Pyrosequencing-based assessment of bacterial community structure along different management types in German forest and grassland soils. PLoS One 6: e17000.
7. Schechtersbach F, Hausmann K, Wylezich C, Weitere M, Arnlt H. (2010) Large-scale patterns in biodiversity of microbial eukaryotes from the abyssal sea floor. Proc Natl Acad Sci USA 107: 115–120.
8. Pointing SB, Chan Y, Lapic DC, Lau MCY, Jungers JA, et al. (2009) Highly specialized microbial diversity in hyper-arid polar desert. Proc Natl Acad Sci USA 106: 19964–19969.
9. Angenent LT, Kelley ST, St Amand A, Pace NR, Hernandez MT. (2005) Metagenomic study of the oral microbiota by Illumina high-throughput sequencing. J Microbiol Methods 79: 266–271.
10. Qin J, Li R, Arumugam M, Burgdorf KS, et al. (2010) A human gut microbial gene catalogue established by metagenomic sequencing. Nature 464: 59–65.
11. Keijer BJ, Zaura E, Huse SM, van der Vossen JMBM, Schuren FHJ, et al. (2008) Pyrosequencing analysis of the oral microflora of healthy adults. J Dent Res 87: 1016–1020.
12. Mobberley JM, Ortega MC, Foster JS. (2012) Comparative microbial diversity analyses of modern marine thromboidic mats by barcoded pyrosequencing. Environ Microbiol 14: 82–100.
13. Furrie E. (2006) A molecular revolution in the study of intestinal microflora. Gut 55: 141–143.
14. Gill SR, Pop M, DeBoy RT, Eckburg PB, Turnbaugh PJ, et al. (2006) Metagenomic analysis of the human distal gut microbiome. Science 312: 1355–1359.
15. Zoetendal EG, Rajilic-Stojanovic M, de Vos WM. (2008) High-throughput diversity and functionality analysis of the gastrointestinal tract microbiota. Gut 57: 1605–1613.
16. Mobberley JM, Ortega MC, Foster JS. (2012) Comparative microbial diversity analyses of modern marine thromboidic mats by barcoded pyrosequencing. Environ Microbiol 14: 82–100.
17. Dusko Ehrlich S, MetaHIT consortium. (2010) Metagenomics of the intestinal microflora: potential applications. Gastroenterol Clin Biol 34 Suppl 1: S23–R.
18. Peleg AY, Hooper DC. (2010) Hospital-acquired infections due to gram-negative bacteria. N Engl J Med 362: 1804–1813.
19. Weber DJ, Raasch R, Rutala WA. (1999) Nosocomial Infections in the ICU. The growing importance of antibiotic-resistant pathogens. Chest 115: 34S–41S.
20. Weber DJ, Raasch R, Rutala WA. (1999) Nosocomial Infections in the ICU. The growing importance of antibiotic-resistant pathogens. Chest 115: 34S–41S.
21. Weber DJ, Raasch R, Rutala WA. (1999) Nosocomial Infections in the ICU. The growing importance of antibiotic-resistant pathogens. Chest 115: 34S–41S.
22. Weber DJ, Raasch R, Rutala WA. (1999) Nosocomial Infections in the ICU. The growing importance of antibiotic-resistant pathogens. Chest 115: 34S–41S.
23. Weber DJ, Raasch R, Rutala WA. (1999) Nosocomial Infections in the ICU. The growing importance of antibiotic-resistant pathogens. Chest 115: 34S–41S.
24. Weber DJ, Raasch R, Rutala WA. (1999) Nosocomial Infections in the ICU. The growing importance of antibiotic-resistant pathogens. Chest 115: 34S–41S.
investigations using the Roche Genome Sequencer FLX platform. BioTechniques 51: 127–133.

25. Nakamura S, Nakaya T, Iida T (2011) Methagenomic analysis of bacterial infections by means of high-throughput DNA sequencing. Exp Biol Med 236: 969–971.

26. Chakravertty S, Helb D, Burday M, Connell N, Alland D (2007) A detailed analysis of 16S ribosomal RNA gene segments for the diagnosis of pathogenic bacteria. J Microbiol Methods 69: 330–339.

27. Baker GC, Smith JJ, Cowan JA (2008) Review and reanalysis of domain-specific 16S primers. J Microbiol Methods 55: 541–555.

28. Kumar PS, Brooker MR, Dowd SE, Camerlengo T (2011) Target region selection is a critical determinant of community fingerprints generated by 16S pyrosequencing. Plos One 6: e20956.

29. Costello EK, Lauber CL, Hamady M, Fierer N, Gordon JI, et al. (2009) Bacterial community variation in human body habitats across space and time. Science 326: 1694–1697.

30. Hanberger H, Walther S, Leone M, Barie PS, Rello J, et al. (2011) Increased mortality associated with meticillin-resistant Staphylococcus aureus (MRSA) infection in the Intensive Care Unit: results from the EPIC II study. Int J Antimicrob Agents 38: 331–335.

31. Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ (1990) Basic local alignment search tool. J Mol Biol 215: 403–410.

32. Cole JR, Wang Q, Cardenas E, Fish J, Chai B, et al. (2009) The Ribosomal Database Project: improved alignments and new tools for rRNA analysis. Nucleic Acids Res 37: D141–D145.

33. Schloss PD, Gevers D, Westcott SL (2011) Reducing the effects of PCR amplification and sequencing artifacts on 16S rRNA-based studies. PLoS One 6: e27310.

34. DeSantis TZ, Hugenholtz P, Larsen N, Rojas M, Brodie EL, et al. (2006) Greengenes, a chimera-checked 16S rRNA gene database and workbench compatible with ARB. Appl Environ Microbiol 72: 5069–5072.

35. Pruesse E, Quast C, Knittel K, Fuchs BM, Ludwig W, et al. (2007) SILVA: a comprehensive online resource for quality checked and aligned ribosomal RNA sequence data compatible with ARB. Nucleic Acids Res 35: 7188–7196.

36. Collins MD, Hoyles L, Hutson RA, Foster G, Falsen E (2001) Corynebacterium testudinoris sp. nov., from a tortoise, and Corynebacterium felinum sp. nov., from a Scottish wild cat. Int J Syst Evol Microbiol 51: 1349–1352.

37. Ciric L, Mullany P, Roberts AP (2011) Antibiotic and antiseptic resistance genes are linked on a novel mobile genetic element: Tn6087. J Antimicrob Chemother 66: 2235–2239.

38. Schloss PD, Gevers D, Westcott SL (2011) Ironing out the wrinkles in the rare biosphere through improved OTU clustering. Environ Microbiol 12: 1889–1898.