Comparative Metagenomic Analysis of Coral Microbial Communities Using a Reference-Independent Approach

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Abstract

By comparing the SEED and Pfam functional profiles of metagenomes of two Brazilian coral species with 29 datasets that are publicly available, we were able to identify some functions, such as protein secretion systems, that are overrepresented in the metagenomes of corals and may play a role in the establishment and maintenance of bacteria-coral associations. However, only a small percentage of the reads of these metagenomes could be annotated by these reference databases, which may lead to a strong bias in the comparative studies. For this reason, we have searched for identical sequences (99% of nucleotide identity) among these metagenomes in order to perform a reference-independent comparative analysis, and we were able to identify groups of microbial communities that may be under similar selective pressures. The identification of sequences shared among the metagenomes was found to be even better for the identification of groups of communities with similar niche requirements than the traditional analysis of functional profiles. This approach is not only helpful for the investigation of similarities between microbial communities with high proportion of unknown reads, but also enables an indirect overview of gene exchange between communities.

Introduction

Metagenomics revolutionized microbial ecology and environmental microbiology because it allows the study of genes and functions of the whole microbial community of an environment, including uncultured micro-organisms. The number of metagenomic studies is growing fast in the last few years, especially due to the rising of ultra-high throughput sequencing technologies. Comparing two or more metagenomics datasets derived from different environments is one common approach used to identify functions and genes that play an important role in one of these environments. Usually, these comparisons are made through functional profiles derived from the annotated sequences according to a reference database, such as SEED, Pfam and COGs. Nevertheless, in several cases, most of the metagenomic sequences cannot be allocated to one of these functional categories due to dissimilarity between newly generated and reference database sequences [1]. Thus, ignoring the portion of unknown sequences during comparative metagenomics can lead to biased conclusions. Some authors have developed comparative analyses of metagenomes using a reference-independent approach. Dutilh et al. [1] proposed the use of cross-assembling to identify similarity relationships among metagenomes. Using cross-assembling files from two different datasets, the crAss method enables the calculation of distances between the metagenomic sample pairs using the number of reads from each metagenome that was used to build a contig. Despite the high efficiency of this method of metagenome comparison, it is computationally intense and time-consuming, especially for requiring assembling of sequences, which is particularly difficult in the case of complex sequence datasets and requires several gigabytes of available RAM memory [2]. Therefore, in this study, a database-independent approach was used to compare different metagenomes (two coral metagenomes sequenced in this work and 29 metagenomes publicly available at MG-RAST) by all-against-all BLAT of the reads (with a threshold of 99% of nucleotide identity and minimum alignment length of 100 bp), in order to find nearly identical sequences that were shared between metagenome pairs. Our hypothesis was that similar microbial communities, with similar functional composition or that harbour the same biological entities (bacterial strains or plasmids, for example) share, together, more identical sequences than distinct communities. The two coral species (Musisamillia hispida and Madracis decactis) utilized in this study are the only scleractinian corals from the São Paulo State coast, and the study of M. hispida is of particular interest because it belongs to a genus endemic to the Brazilian coast.

An advantage of this approach is that the presence of shared identical sequences may also be indicative of recent events involving gene swapping between environments. Using a similar approach, Kloesges et al. [3] analyzed 329 proteobacteria genomes and found that most of the gene sharing was among bacteria from different taxa inhabiting the same habitat. Smillie et al. [4] assessed recently transferred genes (more than 99% nucleotide identity) among 2,235 bacterial genomes, and also
found that the habitat influenced these events, rather than geographical distance or taxonomy. Although there have been several comprehensive studies of gene transference among bacterial genomes, few investigations have considered the way in which genes can move between different environments. Forsberg et al. [5] recently reported multi-drug-resistant soil bacteria containing resistance genes with identical nucleotide sequences to genes from human pathogens, indicative of recent gene exchange events between environmental bacteria and clinical pathogens. However, the authors did not suggest ways in which these genes could be transferred. Thus, the indirect study of the frequency and dynamics of gene swapping events among environments, provided by the approach proposed in this work, may help to understand how genes, such as the ones related to antibiotic resistance, are spread over large-scale geographic distances.

Materials and Methods

Sample collection

One piece of colony (around 5 cm²) of each of the corals Mussismilia hispida and Madracis decactis were collected in March 2012 at Buzios Island, São Paulo State, Brazil (23° 48' 157” S, 45° 07’ 181” W). The sea surface temperature was 22°C, and the coral samples were collected at a depth of 11 m. The colonies were transported to the land in a sterile box with seawater, then briefly rinsed with autoclaved seawater and stored at −20°C for two days prior to DNA isolation.

The specimens were sampled and provided by CEBIMar-USP (Centro de Biologia Marinha da Universidade de São Paulo). The sampling location is a public beach, thus no specific permissions were required to collect the material necessary for the present study.

DNA isolation and sequencing

The coral pieces were vigorously washed with TE buffer (10 mM Tris, 1 mM EDTA, pH 8.0) in a vortex mixer. The solution obtained was centrifuged for 30 s at 8944 RCF units (g) to pelletize the debris. The supernatant was transferred to another tube and centrifuged for 10 min at 15,115 g. The pellet was employed for DNA isolation using a Wizard Genomic DNA Purification Kit (Promega, Madison, Wisconsin, USA), according to the manufacturer’s instructions for Gram-positive bacteria. Average yield was 2 ng/µL of DNA for each sample. The DNA was amplified using a REPLI-g Midi-kit (QIAGEN, Duesseldorf, Germany), according to the manufacturer’s instructions. The DNA was quantified using a Qubit Kit (Invitrogen, Carlsbad, California, USA), and the integrity was confirmed by 1% agarose gel electrophoresis. The DNA obtained ranged between 48,000 and 12,000 bp, and was sequenced by using 454 technology on a Roche GS FLX platform at the DNA Facility of Iowa State University.

Sequence and statistical analysis

Reads were trimmed by quality (minimum of 25) and length (minimum of 100 bp) using PRINSEQ [6]. The barcodes were removed with TagCleaner software [7]. Reads showing 95% of nucleotide identity with sequences of vectors or mitochondrial databases (ftp://ftp.ncbi.nih.gov/refseq/) were removed by using the script exclude_seqs_by_blast.py. The sequences were de-replicated and annotated in SEED subsystems (e-value <1e-05) using the automatic MG-RAST platform [8], and the metagenomes of M. hispida and M. decactis were deposited under the accession numbers 4516694.3 and 4516541.3, respectively. In addition, the de-replicated reads were also submitted to CoMet [9] to identify the Pfam domain families (e-value <0.001) (http://pfam.sanger.ac.uk/). The reads of the metagenomes were also submitted to MetaVir, a web-server for virome analysis [10].

One-way analysis of similarity (ANOSIM) and non-metric multidimensional scaling (MDS), using Bray-Curtis dissimilarity, were performed with PAST software (http://folk.uio.no/ohammer/past/). The differentially abundant functions and Pfam domains were identified using White’s non-parametric t-test with 1,000 bootstrapping replications [11]. The differentially abundant SEED functions where identified with STAMP software (http://kiwi.cs.dal.ca/Software/STAMP). Comparisons between M. hispida and M. decactis libraries were performed using bootstrapping and Bonferroni correction of p-values; Comparison between the coral metagenomes sequenced in this work and the 29 metagenomes publicly available on MG-RAST were also performed in STAMP, using White’s non-parametric t-test and Bonferroni correction of p-values.

In order to detect identical sequences shared among environments, an all-against-all blast (minimum identity 99% and minimum alignment length = 100 bp) was performed with the de-replicated sequences of the metagenomes. A matrix of pairwise Jaccard’s similarities was built using the formula: $J_{AB} = \frac{H_{AB}}{H_{AA} + H_{BB} - H_{AB}}$, where $H_{AB}$ is the number of hits from metagenome A, using metagenome B as the database, $H_{BA}$ is the number of hits from metagenome B, using metagenome A as the database, and $N_A$ and $N_B$ are the total numbers of reads of metagenomes A and B, respectively. Mantel’s test was used to assess the correlation of the Jaccard’s indexes of shared identical sequences with the geographical distance and the Bray-Curtis similarity indexes of taxonomic (at Class level) and functional composition (at SEED level 2, 3, and Pfam domains), using PAST software. A gene-sharing network was visualized using Cytoscape v. 3.0 software, and the metagenomes were represented by the nodes and the edges showing Jaccard’s similarity values equal to or greater than 1.0.$10^{-6}$. A network was produced using a prefrontal force directed algorithm, weighted with Jaccard’s values. Nodes were clustered using the ModuLand plug-in [12], which searches for groups with a high density of internal edges, weighted with the Jaccard’s values.

Results

Description of the metagenome of the Brazilian corals

A total of 368,772 good quality reads were obtained for Mussismilia hispida, with an average length of 446 bp, and 293,580 for Madracis decactis, with an average length of 453 bp. Only 4.4% (16,910) of the reads from M. hispida and 3.3% (9,688) of the reads from M. decactis could be annotated by the hierarchical SEED subsystem, and 8.2% (30,239) of the reads from M. hispida and 8.2% (24,073) of M. decactis presented Pfam hits. Table 1 shows the top ten Pfam domains and SEED functions identified for each coral species; a high abundance of phage and virus proteins can be seen for both classification systems and libraries. The features (at Level 2) that were different for the two metagenomes and presented the lowest p-values and the highest similarity indexes of taxonomic (at SEED level 2, 3, and Pfam domains), using PAST software. A gene-sharing network was visualized using Cytoscape v. 3.0 software, and the metagenomes were represented by the nodes and the edges showing Jaccard’s similarity values equal to or greater than 1.0.$10^{-6}$. A network was produced using a prefrontal force directed algorithm, weighted with Jaccard’s values. Nodes were clustered using the ModuLand plug-in [12], which searches for groups with a high density of internal edges, weighted with the Jaccard’s values.
Pseudomonadaceae, corresponding to 25.1% (4,290) of the bacterial reads in *M. hispida* and 21.6% (4,052) of the bacterial reads in *M. decactis*. Conversely, according to CoMet Pfam classification, most of the reads, for both corals, were classified as VMG (Viral Metagenome) (Table 2). According to the MetaVir taxonomic classification, 30,619 (10.4%) reads of the *M. decactis* metagenome were of virus origin, and 78% (23,872) of these reads were classified as ssDNA viruses, with the Circoviridae family being the most representative, with 31% of the viral reads (9,392). In the metagenome of *M. hispida*, 34,160 (9.2%) reads were from viruses, with 80% (27,191) classified as ssDNA viruses and the most representative family being the Microviridae/Gokushovirinae, with 29% (9,829) of the viral reads.

Comparative metagenomics

The Pfam and SEED function distributions of the *M. hispida* and *M. decactis* metagenomes were compared with 29 metagenomes publicly available at MG-RAST, including the metagenomes of two coral species, human gut, chicken cecum, invertebrates, seawater, soil, mines, plasmidomes of cow rumen and activated sludge, and viromes of cystic fibrosis lung and seawater (Table S1).

The distribution of Pfam domains in the coral metagenomes was significantly different from the other metagenomes (one-way ANOSIM, R = 0.8277, p = 0.0069). Those overrepresented in the Brazilian coral metagenomes were: PF02674 (difference around 108-fold, p = 0.0001), related to colicin V production in *E. coli*; PF02892, a BED-type zinc finger domain related to eukaryotic transposases (difference around 10-fold, p = 0.0009); PF03239 (difference around 33-fold, p = 0.0002), described as an iron permease FTR1 family; PF07903, described as PaRep2a protein of unknown function (difference around 2-fold, p = 0.0006); PF08668, involved in nucleic acid metabolism and signal transduction (difference around 3-fold, p = 0.0005); and PF11654, a domain of unknown function that seems to be involved in protein export (difference around 316-fold, p < 0.0001).

At SEED Level 2, the differences between the present data and publicly available datasets were significant (one-way ANOSIM, Figure 1. Distribution of the SEED categories (Level 2) with difference between the metagenomes of *M. hispida* (light gray) and *M. decactis* (dark gray).

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### Table 1. Functional classification of *M. hispida* and *M. decactis* reads performed using MG-RAST and CoMet platforms.

| Mussismilia hispida | Madracis decactis |
|---------------------|-------------------|
| Pfam                | Pfam Description  | % of total reads | Pfam | Pfam Description  | % of total reads |
| PF02305 Capsid protein (F protein) | 1.8659 | PF02407 Putative viral replication protein | 1.6346 |
| PF00910 RNA helicase | 0.8976 | PF00910 RNA helicase | 1.1833 |
| PF02407 Putative viral replication protein | 0.6847 | PF01446 Replication protein | 0.7610 |
| PF00124 Photosynthetic reaction centre protein | 0.5396 | PF06280 Fn3-like domain (DUF1034) | 0.5338 |
| PF01844 HNH endonuclease | 0.2386 | PF01844 HNH endonuclease | 0.4881 |
| PF01312 Replication protein | 0.2251 | PF01312 Replication protein | 0.7610 |
| PF09295 ChiPs (Chs5p-Af1p-binding proteins) | 0.1908 | PF08019 Domain of unknown function (DUF1034) | 0.3546 |
| PF01051 Initiator replication protein | 0.1906 | PF02305 Capsid protein (F protein) | 0.3406 |
| PF06280 Fn3-like domain (DUF1034) | 0.1003 | PF04127 DNA/pantothenate metabolism flavoprotein | 0.2742 |
| PF00006 ATP synthase alpha/beta family | 0.0998 | PF00006 Geminivirus Rep catalytic domain | 0.2153 |
| SEED Function | % of total reads | SEED Function | % of total reads |
| Phage protein | 0.01247 | Phage protein | 0.0361 |
| Gamma-carotene hydroxylase | 0.00597 | Ribonucleotide reductase of class la (aerobic), alpha subunit (EC 1.17.4.1) | 0.0061 |
| Heat shock protein 60 family chaperone GroEL | 0.00569 | Gamma-carotene hydroxylase | 0.0061 |
| COG0009 Sua5 subfamily, required for N6-threonylcarbamoyl adenosine t(6)A37 modification in tRNA | 0.00569 | Heat shock protein 60 family chaperone GroEL | 0.0051 |
| Ribonucleotide reductase of class la (aerobic), alpha subunit (EC 1.17.4.1) | 0.00542 | COG0009 Sua5 subfamily, required for N6-threonylcarbamoyl adenosine t(6)A37 modification in tRNA | 0.0051 |
| ATP synthase beta chain (EC 3.6.3.14) | 0.00515 | ATP-binding protein | 0.0044 |
| CoB—CoM heterodisulfide reductase subunit C (EC 1.8.98.1) | 0.00515 | Nitrile reductase probable electron transfer Fe-S subunit (EC 1.7.1.4) | 0.0044 |
| Photosystem II protein D2 (PsbD) | 0.00461 | Decarboxylase | 0.0041 |
| Xaa-Pro aminopeptidase (EC 3.4.11.9) | 0.00380 | Translation elongation factor LepA | 0.0031 |
| Type I restriction-modification system, specificity subunit S (EC 3.12.1.3) | 0.00380 | GMP synthase [glutamine-hydrolyzing] (EC 6.3.5.2) | 0.0027 |

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At SEED Level 3, the metagenomes of *M. hispida* and *M. decactis* were also more similar to each other than to the other metagenomes (one-way ANOSIM, \( R = 0.5653, \ p = 0.0328 \)). The functional categories at SEED Level 3 that showed significant differences between the metagenomes of the Brazilian corals and all the other metagenomes were identified using White's non-parametric t-test with 1,000 bootstrapping replications. Figure 2 shows the attributes where the differences presented the smallest p-value. The only SEED categories that were differentially distributed were “Accession colonization factor”, “Dot-Icm type IV secretion system”, and “Type III secretion systems”, which were more abundant in the metagenomes of the Brazilian corals.

The visualization of similarities among the metagenomes by non-metric MDS of SEED functional categories and Pfam profiles did not result in clear discriminations of the communities (Figures S1 and S2).

No Pfam domains or SEED categories were found to be unique to the coral metagenomes used in this study (*M. hispida*, *M. decactis*, *Porites compressa* and *Acropora* sp.).

**Gene-sharing network**

An all-against-all approach was used to look for nearly identical sequences (99% of nucleotide identity and minimum alignment length of 100 bp) in pairs of metagenomes, in order to obtain their similarity indexes (Table S2). The metagenomes that shared more identical sequences were *M. decactis* and *M. hispida* (\( J = 9.152 \times 10^{-4} \)). The human gut metagenome TS5 shared more identical sequences with the chicken cecum sample (\( J = 3.47 \times 10^{-2} \)) than with the human gut TS1 (\( J = 1.53 \times 10^{-4} \)). Other metagenomes that shared a high number of identical sequences were the marine sponges SpongeAb1 and SpongeAb2 (\( J = 4.21 \times 10^{-4} \)), and the artic viromes ArcticVir and GOMVir (\( J = 3.13 \times 10^{-4} \)). Pfam domains were identified in the shared sequences of the four pairs of metagenomes that most shared sequences (*M. hispida* and *M. decactis*; ArcticVir and GOMVir; SpongeAb1 and SpongeAb2; and Gut_Ts5 and ChickenCecum). The top ten Pfam hits are shown in Table S3. However, on average, Pfam domains were identified in only 26% of the sequences. The Jaccard’s values of shared sequences were not strongly correlated with the taxonomic profile at superkingdom level (Mantel’s test, \( R = 0.1983, \ p = 0.0002 \)), the SEED functional profile at level 3 (Mantel’s test, \( R = 0.1172, \ p = 0.0028 \)), and the Pfam distribution (Mantel’s test, \( R = 0.1892, \ p = 0.0002 \)). This means that the metagenome pairs that were more similar in terms of functional and taxonomic (at the superkingdom level) composition did not share more identical sequences. Moreover, no strong correlation was found between sequence-sharing and geographical distance (Mantel’s test, \( R = 0.1638, \ p = 0.0002 \)), hence a metagenome pair collected at close sample sites did not share more sequences than a metagenome pair collected at distant sample sites. ANOSIM was used to test whether datasets for groups in the same category shared more sequences within the group than between groups. The categories tested were habitat (marine or terrestrial); lifestyle (free-living or animal-associated); climate (tropical, temperate, or polar); metagenome type (microbial, plasmidial, or viral); sequencing methodology (cloning or not); MDA (amplified or not); and study (metagenomes sequenced in the same study were grouped together) (Table 3). The groups that had significant differences in sequence-sharing were marine and terrestrial (ANOSIM, \( R = 0.2032, \ p = 0.0003 \)), and tropical and temperate (ANOSIM, \( R = 0.2109, \ p = 0.0001 \)).

A visualization of the similarities among the metagenomes obtained by non-metric MDS of gene-shared hits is presented in

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**Table 2**

| CoMet (TaxyPro) Taxonomic classification (%) | MG-RAST Taxonomic classification (%) | VMG = viral metagenomes |
|---|---|---|
| Archaea | Bacteria | Eukaryota | Archaea | Bacteria | Eukaryota | Viruses | Archaea | Bacteria | Eukaryota | Viruses |
| *Mussismilia hispida* | 1.39 | 57.76 | 29.31 | 0.43 | 19.2 | 7.1 | 6.4 | 5.9 | 8.3 |
| *Madracis decactis* | 0.89 | 65.26 | 15.30 | 5.77 | 11.5 | 7.1 | 6.4 | 5.9 | 8.3 |

VMG = viral metagenomes.

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**Reference**

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Figure 2. Distribution of the SEED categories (Level 3) with difference between the metagenomes of *M. hispida* and *M. decactis* (light gray) and the 29 metagenomes publicly available at MG-RAST (dark gray).
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Figure S3, for comparison with the functional similarities ( Pfam and SEED profiles) using the same technique.

Figure 3 shows the gene-sharing network among metagenomes, where thicker edges represent greater Jaccard’s values between two metagenomes (nodes). Table S4 shows the values of centrality measures of each node. Polynesia metagenome presented the highest values for degree and betweenness centrality, followed by Shudge_V09 and WaterjF1 (Table S4).

Nine clusters were detected among the metagenomes analyzed. A cluster is a highly dense region in a network, which means that clusters are groups of nodes that share more identical nucleotide sequences. Nodes belonging to the same module are presented in the same colour in Figure 3. For example, *M. hispida*, *M. decactis*, and SARVir belong to the same module, which means that they shared more sequences with each other than with other nodes.

Discussion

Although very similar, the functional and taxonomic profiles of *M. hispida* and *M. decactis* presented some unique features. Notable were the differences in the dominant viral family. The Circoviridae was the most abundant in the *M. decactis* dataset. Members of this family are usually described as infecting animal cells [13]. The Microviridae family, most abundant in *M. hispida*, infects bacteria [14]. Further studies, with a greater number of coral colonies collected on different days and during different seasons, are needed to better understand the structure and specificity of the viral communities of these coral species. However, several sequences of the Gokushovirinae group were found in both libraries. Gokushoviruses belong to the Microviridae family and have been reported in several metagenomic studies, but they were never isolated and their hosts remain unknown [15]. Gokushovirus sequences comprised nearly 6% of the reads of the Sargasso Sea Virome (SARVir) [16], which can explain the similarities found between the *M. hispida*, *M. decactis*, and SARVir datasets. Labonte & Suttle [15] suggested that these viruses are highly virulent and can become dominant during a lytic event, because they have small genomes (around 4–5 kb) that permit rapid replication.

The way in which coral-bacteria associations are established is one of the main concerns of coral microbiology studies. Marine bacteria colonize the coral after larval settlement [17]. The assembly of microbial communities associated with marine organisms seems to be a process that is partly selective and partly random [18]. It is already well known that the microbial communities of corals are different from those of the surrounding environment [19,20,21], but there is little evidence concerning the functions required for the establishment and maintenance of a bacteria-coral association [22,23,24]. Comparative studies between coral microbial communities and other environments brings some insights into genes that could be advantageous for the coral niche. Comparison of the functional profiles of the present metagenomes with other metagenomes enabled the observation of some interesting features of the metagenomes of Brazilian corals, such as the relative enrichment of genes related to a colicin V production protein (CvpA), PF02674. The plasmid-borne cvpA gene encodes an inner membrane channel required for the production/secretion of the colicin V toxin in *E. coli* [25]. It has also been found in an aphid symbiont [26] and endosymbionts of deep-sea tubeworms [27]. *E. coli* cvpA mutants presented alterations in biofilm properties and curli fibers [28]. The glycosyltransferase-like family 2 (PF1011) includes putative proteins involved in capsule biosynthesis [29]. Glycosyltransferases are proteins responsible for the synthesis of polysaccharides or glycoproteins [30], and are related to the symbiotic colonization of *V. fischeri* [31], which has been suggested to be laterally transferred and advantageous in host gut [32]. Thus, the CvpA protein and glycosyltransferases are apparently involved in symbiotic relationships, and might act to enhance bacterial colonization of corals.

Comparisons of SEED functional profiles also revealed important functions in the metagenomes of the Brazilian corals. Genes that encode proteins involved in protein secretion systems type III (T3SS) and type IV (T4SS) were found to be enriched in our metagenomes. Protein secretion systems play a central role in modulating bacteria-host interactions. T3SS are found in pathogens and commensal bacteria that interact with both plant and animal hosts [33]. Several T3SS genes are encoded in pathogenicity islands or are located on plasmids, and are commonly subject to horizontal gene transfer [34]. T3SS has been frequently found in arthropoda-associated microbial communities [35]. Bondarev et al. [36] suggested that T3SS is involved in *Pseudovibrio* interactions with marine invertebrates. T4SS is able to transport not only protein but also DNA [33]. This system has been found enriched in microbial communities where HGT events are advantageous, because of intense selective pressure, and is often encoded on self-transmissible plasmids and integrative conjugative elements [33]. Based on this information, we suggest that these protein secretion systems can play a role in the establishment and tightening of bacterial interaction with the coral host, and for this reason they are subject to HGT and were found to be enhanced in the Brazilian coral metagenome.

In this study, most of the metagenome pairs shared relatively few identical sequences (254 of 463 comparisons resulted in J < 1.0 x 10^-6). The dataset pairs with highest J values included some metagenomes belonging to the same project and prepared by the same group, such as ArcticVir and GOMVir [16], SpongeAb1 and SpongeAb2 [37], and the two metagenomes sequenced in this work. In contrast, the metagenomes Gut_TSS [38] and ChickenCecum [39] were sequenced by different groups. Given its nature, the analysis presented here may be influenced by the methodologies employed in different studies. However, we have...
found that metagenomes with the same sequencing methodology (cloning) or amplification methodology (MDA) do not share more sequences. Despite the limitations of the present results, some conclusions can be drawn that might help to guide future studies of identical nucleotide sequences that are shared between metagenomes. Network and node properties obtained from bacterial metagenome gene-sharing data can help to identify critical environments and factors for the swapping of genes among microbial communities. The betweenness centrality measure is defined as the frequency with which a node lies on the shortest path between two other network nodes, and a metagenome with a high betweenness value can transfer sequences to many other metagenomes in the network, with a low number of gene transfer events. In other words, it can function as a bridge between disconnected regions of the network [40]. Our results indicated that marine microbial communities may play this role. In epidemiologic terms, the marine environment presents higher degree of connectivity than the terrestrial one, and one of the reasons is the lack of dispersal barriers [41]. Therefore, microorganisms such as bacteria and viruses can spread easily and rapidly through oceans, which may explain the high betweenness values found in the Polynesia and WaterJF1 metagenomes. Viruses may be one of the most important carriers of genes through biomes, due to their ubiquity and abundance [42,43]. Corroborating this notion, two studies have found a worldwide distribution of identical viral sequences [44,45], showing that viruses can move between different environments.

Cluster or modularity analysis allows the identification of groups of nodes that share more sequences between them than with nodes outside the group. These groups can represent microbial communities with similar taxonomic or functional composition, or groups of communities where gene swapping is more likely to happen. In both cases, despite the impossibility of functional classification of the shared sequences, the metagenome clusters can be interpreted as groups of microbial communities that are under similar selective pressures. Examples are the cluster composed of microbial communities of the vertebrate digestive system (Gut_TS1, Gut_TS5, CowRumen, and ChickenCecum) and the one composed of marine viromes (GOMVir, ArcticVir, BBCVir, and TampaBay). The metagenomes primarily consisting of Atlantic Ocean phage sequences compose another cluster (Mussismilia, Madracis, and SARVir). It is interesting to note that the four coral metagenomes used in this study were not clustered together. One possible explanation for this is the fact that most of coral-derived microbial metagenomes also present host, algae, chloroplast and mitochondrial DNA, which make comparative analyses difficult [24].

Regarding the factors that influence the observed profile of shared sequences among datasets, we have found that climate and habitat can influence the determination of metagenome groups; however, these influences seem to be slight, and other factors not considered in our analysis may be more important, such as physical-chemical and nutritional parameters, ocean and wind currents, and host biogeography, amongst others.

Finally, we consider that comparative metagenomics using a reference-independent approach is even better than using functional profiles (SEED and Pfam), because it led to a more clear discrimination of the microbial communities used in this study (Figures S1, S2 and S3). One explanation for the high overlapping among the datasets observed with SEED and Pfam profiles is that using reference-dependent approaches to compare metagenomes only takes in consideration the small amount of sequences that are similar to known sequences, and ignores the high amount of sequences from unknown organisms and with
unknown function, which may be holding the differences among microbial communities.

Conclusions

Comparative metagenomics allows the identification of microbial communities that share similar functional and taxonomic composition, and the observation of patterns controlling ecological niche partitioning. By comparing the metagenomes associated with two coral species with other metagenomes, we were able to identify functions that are probably either required or advantageous for coral colonization, such as transport proteins.

Besides of the simplicity of the reference-independent approach suggested in this work, the analysis of identical sequences shared among the microbial communities was found to be even better for the identification of groups of communities with similar niche requirements than the traditional analysis of functional profiles. This approach is not only helpful for the investigation of similarities between microbial communities, but also enables an indirect overview of gene exchange between communities. We consider this to be a successful procedure for use in comparative metagenomics, and we also suggest the use of tblastx comparisons in future studies.

Supporting Information

Figure S1 Visualization of Bray-Curtis similarities of Pfam profiles between metagenomes using non-metric multidimensional scaling. Stress value = 0.1048. (TIF)
Figure S2 Visualization of Bray-Curtis similarities of Level 3 (SEED subsystem) profiles between metagenomes using non-metric multidimensional scaling. Stress value = 0.1868. (TIF)

Figure S3 Visualization of Bray-Curtis similarities of gene-shared hits between metagenomes using non-metric multidimensional scaling. Stress value = 0.2484. (TIF)

Table S1 Information about the metagenomes utilized in this work. (DOCX)

Table S2 Jaccard’s similarities among the metagenomes obtained from gene-sharing data. (DOCX)

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