Exploring the interaction patterns in seasonal marine microbial communities with network analysis

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Abstract—With the development of high-throughput and low-cost sequencing technology, a large amount of marine microbial sequences is generated. So, it is possible to research more uncultivated marine microbes. The interaction patterns of marine microbial species and marine microbial diversity are hidden in these large amount sequences. Understanding the interaction pattern and structure of marine microbe have a high potential for exploiting the marine resources. Yet, very few marine microbial interaction patterns are well characterized even with the weight of research effort presently devoted to this field. In this paper, based on the 16S rRNA tag pyrosequencing data taken monthly over 6 years at a temperate marine coastal sits in West English Channel, we employed the CROP unsupervised probabilistic Bayesian clustering algorithm to generate the operational taxonomic units (OTUs), and utilized the PCA-CMI algorithm to construct the spring, summer, fall, and winter seasonal marine microbial interaction networks. From the four seasonal microbial networks, we introduced a novel module detecting algorithm called as DIDE, by integrating the dense subgraph, edge clustering coefficient and local modularity, to detect the interaction pattern of marine microbe in four seasons. The analysis of network topological parameters shows that the four seasonal marine microbial interaction networks have characters of complex networks, and the topological structure difference among the four networks maybe caused by the seasonal environmental factors. The marine microbial interaction patterns detected by DIDE algorithm in four seasons show evidence of seasonally interaction pattern diversity. The interaction pattern diversity of fall and winter is more than that of spring and fall, which indicates that the seasonal variability might have the greatest influence on the marine microbe diversity.

Keywords—marine microbe; operational taxonomic unit (OTU); interaction pattern; network; clustering

I. INTRODUCTION

Marine microbes are the ‘gatekeepers’ for the Earth System with an estimated contribution to global primary productive of between 50% and 90% [1], and also are the important composition in the marine ecosystem. Yet specific ecological relationships among these taxa are largely unknown. This is part due to the dilute, microscopic nature of the planktonic microbial community, which prevents direct observation of their interactions [2]. With the development of high-throughput DNA sequencing technologies that yield a mass of reads of small-subunit rRNA gene (16S rRNA/18S rRNA) and DNA, we can describe the compositions of microbial communities, their diversity and how communities may change across space, time or experimental treatments based on these sequence data [3]. However, most of the current analytical approaches of describing and comparing the structure of communities often focus on the total numbers of taxa, the relative abundances of individual taxa and the extent of phylogenetic or taxonomic overlap between communities or community categories[4-6]. In contrast, there has been far less attention focused on using sequence data to explore the direct or indirect interactions between microbial taxa coexisting in environmental samples. Although some researchers used the network analysis to explore co-occurrence pattern in soil and ocean[2, 3, 7-9], they just constructed the associate networks to show the co-occurrence pattern, and did not further mine the networks finding the pattern structures. The microbial interaction (or co-occurrence) patterns can offer new insight into the structure of complex microbial communities, reveal the niche spaces shared by community members, identify habitat affinities or shared physiologies that could guide more experimental settings. In this article, we will construct the spring, summer, fall and winter marine microbial interaction networks, introducing a new module detecting algorithm to find the microbial interaction patterns. The aim was to understand the relationship among microbe and seasonal variability and try to determine the microbial interaction pattern difference among seasons.

II. MATERIAL AND METHODS

A. Dataset

The 16S rRNA sequence dataset used in this paper were downloaded from http://vamps.mbl.edu/index.php, which includes 969,400 sequences generated from 76 time point seawater samples at the surface of L4 sampling site in the West English Channel [8]. The 76 seawater samples were arranged into spring (March-May), summer (June-August), fall (September-November) and winter (December-January) seasons, in which 22, 27, 13 and 14 samples belong to spring, summer, fall and winter seasons respectively. And the 16S rRNA sequence numbers of spring, summer, fall and winter seasons are 249,395, 293,549, 202,356 and 224,100, respectively. In order to establish the seasonal marine microbial network at the taxonomic level (e.g. species, genus), the 16S
rRNA sequences were grouped into species-level operational taxonomic units (OTUs) with CROP program [10], which resulted in 5858 OTUs. CROP adopts an unsupervised probabilistic Bayesian clustering algorithm and uses a soft threshold for defining the OTUs, which bypasses setting a subjective hard cut-off threshold, thus may effectively reduce the effects of PCR and sequencing errors in inferring OTUs.

B. OTU-OTU correlation network modeling

In order to research the correlation among different microbial species and explore their interaction patterns, we use a vector \( X_i \) to represent each OTU in the four seasons.

\[
X_i = [x_{i1}, x_{i2}, \ldots, x_{in}, \ldots, x_{is}] \quad (i=1, \ldots, 5858) \tag{1}
\]

where \( x_{is} \) is the \( i \)-th OTU abundance value in the \( s \)-th sampling, that is, \( x_{is} \) equals the ratio of the sequence number \( N_{is} \) contained in the \( i \)-th OTU and the total sequence number \( N_s \) contained in the \( s \)-th sampling. To reduce the sequencing effort bias, the \( x_{is} \) value was set to zero if \( N_s < 5 \). For reducing the false higher correlation between vectors, we also remove these OTU vectors which contain less than 3 non-zero elements. After these processing, we can obtain 825 OTU vectors, in which spring season contains 205, summer 179, fall 208 and winter 233 OTUs respectively. Then, the four microbial abundance matrices of spring, autumn, fall and winter season were constructed by normalizing every OTU vector.

The path consistency algorithm based on conditional mutual information (PCA-CMI) was firstly used to infer (or reconstruct) the gene regulatory networks (GRNs) from gene expression data [11]. In PCA-CMI algorithm, the conditional independence between a pair of genes is represented by conditional mutual information between this gene-pair given certain other genes. PCA-CMI can cover nonlinear relations between gene pairs based on mutual information (MI) and CMI from information theory, and detect the nonlinear statistical dependence. If the seasonal microbial abundance matrices were considered as the gene expression data, we can use PCA-CMI to construct the seasonal marine microbial correlation networks. The process of PCA-CMI can be described simply as follows.

Firstly, generate a complete graph \( G \) according to the number of OTUs. Secondly, compute the zero-order MI \( I(i,j) \) for all OTU pairs.

\[
I(X,Y) = - \sum_{x \in X, y \in Y} p(x,y) \log \frac{p(x,y)}{p(x)p(y)} \tag{2}
\]

If the OTU\(_i\) and OTU\(_j\) has low or zero MI, then the edge between OTU\(_i\) and OTU\(_j\) is deleted. Thirdly, for adjacent OTU\(_i\) and OTU\(_j\), select the adjacent OTU\(_k\) of them and compute first-order CMI \( I(i,j,k) \).

\[
I(X,Y|Z) = - \sum_{x \in X, y \in Y, z \in Z} p(x,y,z) \log \frac{p(x,y|z)}{p(x|z)p(y|z)} \tag{3}
\]

If the OTU\(_i\) and OTU\(_j\) has low or zero CMI which represents independent correlation, then the edge between them is deleted. The next step is to compute higher order CMI until there are no more adjacent edges. In this paper, considering the computational complexity, we just compute the second-order CMI and construct the four second-order marine microbial correlation networks for spring, summer, fall and winter seasons.

C. DIDE network module detecting algorithm

Before introducing the DIDE algorithm, we will present some notions. 1) Number of shared adjacent nodes \( |N_u| \). For an undirected graph \( G(V,E) \), suppose \( N_i \) and \( N_j \) are the adjacent node sets of node \( v_i \) and \( v_j \) respectively, the number of shared adjacent nodes between any two nodes is defined as \( |N_i | = |N_i \cap N_j | \).

2) Dense subgraph. Suppose the degree of node \( v_u \) is maximum, \( N = \{v_1, v_2, \ldots, v_s\} \) is adjacent node set of \( v_u \), \( N' = \{v_1, v_2, \ldots, v_s\} \) is adjacent node set of \( v_u \) shared the most nodes with \( N, N' = \{v_1, v_2, \ldots, v_s\} \) is the adjacent node set of \( v_u \) shared the second most nodes with \( N \), then the dense subgraph is defined as \( C_{ij} = \{v_i \in \{v_1, v_2, \ldots, v_s\} \} \).

3) Edge clustering coefficient \( C_{ij} \). Suppose that there is an edge \( e_{ij} \) between node \( v_i \) and \( v_j \), and the two nodes have a shared node \( v_k \), and the edges \( e_{ik}, e_{jk} \) and \( e_{ij} \) can form a triangle loop, then the edge clustering coefficient \( C_{ij} \) of edge \( e_{ij} \) is defined as \( C_{ij} = (t_i + 1)/\min(d_i, d_j, d_{ij}) \). \( d_i \) and \( d_j \) are the degree of node \( i \) and \( j \) respectively, \( t_i \) is the total number of true triangle loops including the edge \( e_{ij} \), and the denominator represents the total number of theoretical triangle loop including the edge \( e_{ij} \).

The DIDE algorithm can be described in detail as follows.

i) Select the maximum degree node forming the initial dense subgraph \( C_{ab} \), then producing the module \( M \).

ii) Compute the module \( M \) local modularity value \( Q_M = L_{in}/(L_{in} + L_{out}) \), \( L_{in} \) is the inner edge number of the module \( M \), \( L_{out} \) is the outer edge number of the module \( M \).

iii) Search the adjacent node \( v_i \) of the module \( M \); compute the connection tightness \( U_i \) between node \( v_i \) with the module \( M \), \( U_i = |E_{in}|/d_i \). \( |E_{in}| \) means the connection edge number of node \( v_i \) with the module \( M \), \( d_i \) is the degree of node \( v_i \).

iv) If \( U_i \geq \theta \), then add the node \( v_i \) to the module \( M \), forming a new module \( M' \).

v) Else, compute the edge clustering coefficient of other residue adjacent node \( v_i \). If the edge clustering coefficient of node \( v_i \) with the module \( M \) is the biggest, then add the node \( v_i \) to the module \( M \), forming a new module \( M' \).

vi) Compute the \( M' \) local modularity value \( Q_{M'} \). If \( Q_{M'} - Q_M < 0 \), then remove the node \( v_i \) from \( M' \).

vii) Repeat iii) to vii) for all other adjacent node of the \( M' \).
until $\Delta Q_M = 0$.

viii) If there is still other dense subgraph, return to i).

The flow diagram of DIDE algorithm is displayed in Fig.1.

III. RESULTS AND DISCUSSION

A. Topology analysis of four seasonal marine microbial correlation networks

In order to study and analyze the microbial diversity of spring, summer, fall and winter seasons, we constructed the four seasonal marine microbial correlation networks with PCA-CMI algorithm. Fig.2 shows the four marine microbial correlation networks in spring, summer, fall and winter seasons. We also computed their topological parameters including the average degree, average clustering coefficient, average power law degree, modularity, and compared with their corresponding random networks. The comparison results of four seasonal networks and random networks are summarized in Table 1.

Fig. 2. Marine microbial correlation networks in spring, summer, fall and winter seasons.
From Table 1, we can see that there is some difference on the topological parameters among the spring, summer, fall and winter seasonal microbial correlation networks, which maybe caused by the seasonal environmental factors. Compared with random networks, four seasonal microbial correlation networks have bigger average clustering coefficient, average power law degree and modularity, which indicate that the four seasonal microbial correlation networks have some characters of complex network.

B. The interaction patterns in seasonal microbial networks detected by DIDE Algorithm

We firstly choose the typical American College Football network to verify the performance of DIDE algorithm. American Football Network contains 115 nodes and 613 edges [12]. Each node represents one team which belongs to one of the 12 Unions. Edge indicates the games played by the teams against other during the regular season of fall 2000. The performance of DIDE is better than that of Zhang’s method [13], FN[14] and GN [15]. DIDE algorithm with $U_i=0.5$ can detect 12 communities, and the accuracy arrives at 91%. Yet Zhang’s method, FN and GN just detect 10, 10, and 11 communities respectively, and their corresponding accuracies are 90%, 65% and 78% respectively. We also investigate the parameter $U_i$ how to affect the results for four seasonal marine microbial OTU-OTU networks. The numbers of microbial interaction pattern detected by DIDE algorithm for spring, summer, fall and winter microbial OTU-OTU networks are listed in Table 2, and the structures of pattern at $U_i=0.5$ are show in Fig. 3.

The results in Table 2 show that the interaction pattern diversity of fall and winter is more than that of spring and fall, which indicates that the seasonal variability might have the greatest influence on the marine microbe diversity, and the parameter $U_i$ has little influence to the community number detected for $U_i>0.5$.

| TABLE 1. TOPOLOGICAL PARAMETERS OF FOUR SEASONAL MARINE MICROBIAL CORRELATION NETWORKS AND THE CORRESPONDING RANDOM NETWORKS |
|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|
| Node Number | Spring | Summer | Fall | Winter | 1 | 2 | 3 | 4 |
| Edge Number | 1419 | 512 | 1690 | 1791 | 1419 | 512 | 1690 | 1799 |
| Avg. degree | 13.8 | 5.8 | 16.3 | 15.4 | 13.8 | 5.8 | 16.3 | 15.4 |
| Avg. clustering coefficient | 0.37 | 0.20 | 0.43 | 0.343 | 0.065 | 0.015 | 0.073 | 0.065 |
| Avg. power law degree | 0.637 | 0.948 | 0.322 | 0.882 | 0.355 | 0.507 | 0.06 | 0.245 |
| Modularity | 0.461 | 0.571 | 0.553 | 0.493 | 0.223 | 0.377 | 0.212 | 0.219 |

| TABLE 2. NUMBERS OF MICROBIAL INTERACTION PATTERN DETECTED WITH DIFFERENT PARAMETER $U_i$ |
|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|
| 0.4 | 0.5 | 0.6 | 0.7 | 0.8 | 0.9 | 1.0 |
| spring | 4 | 4 | 5 | 6 | 6 | 6 | 6 |
| summer | 2 | 3 | 3 | 2 | 2 | 2 | 2 |
| fall | 10 | 13 | 15 | 15 | 18 | 18 | 17 |
| winter | 1 | 12 | 13 | 18 | 20 | 20 | 20 |

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Fig. 3 shows that the interaction pattern diversity of fall and winter is more than that of spring and fall. According to the annotation information of OTUs at taxonomic level by using a number of different annotation strategies (e.g. GAST[4], BLAST against Greengenes[16], SIVA[17], RDP[18]), we analyzed in detail the biggest module for every seasonal network. The M1 module in spring microbial network has 71 OTUs in which the 44 OTUs were identified in Phylum level as ‘Proteobacteria’, 7 OTUs as ‘Actinobacteria’, 5 OTUs as ‘Bacteroidetes’, 5 OTUs as ‘Deferribacteres’, 4 OTUs as Chloroplast, 1 OTU as ‘Gemmatimonadetes’, 1 OTU as ‘Verrucomicrobia’ and 4 OTUs could not be identified; in family level, the 7 OTUs were identified as ‘Rhodospirillaceae’, 6 OTUs as ‘Rhodobacteraceae’, 5 OTUs as ‘SAR11’, 5 OTUs as ‘SAR406’, 4 OTUs as ‘SAR86’, 3 OTUs as ‘Cryomorphaceae’, 2 OTUs as ‘Comamonadaceae’, 2 OTUs as ‘Eubacteriaceae’, the other OTUs belong to ‘Flammeovirgaceae’, ‘Falvobacteriaceae’, ‘Oceanospirillaceae’, ‘Piscirickettsiaceae’, ‘Propionibacteriaceae’, ‘Pseudomonadaceae’, ‘Salinisphaeraceae’, ‘Sinobacteraceae’, ‘Sphingomonadaceae’, ‘Subdivision3’ respectively, and 22 OTUs could not identified. Due to more 70% OTTUs of M1 in spring network could not be identified to genus, we did not further analyze M1 structure under genus taxonomic level.

The M1 module in summer microbial network has 19 OTUs in which the 12 OTUs in Phylum level belong to ‘Proteobacteria’, 3 OTUs to ‘Bacteroidetes’, 2 OTUs to ‘Deferribacteres’, the other OTUs belong to ‘Cyanobacteria’ and ‘Verrucomicrobia’ respectively.

The M1 module in fall microbial network has 26 OTUs in which the 12 OTUs in Phylum level belong to ‘Bacteroidetes’, 9 OTUs to ‘Proteobacteria’, 3 OTUs to
The M1 module in winter microbial network has 54 OTUs in which the 38 OTUs in Phylum level belong to ‘Proteobacteria’, 10 OTUs to ‘Bacteroidetes’, 2 OTUs to ‘Actinobacteria’, the other two OTUs belong to ‘Cyanobacteria’ respectively.

The M1 structural analysis in four seasonal microbial networks show that a large fraction microbial interactions in Phylum level occur among ‘Proteobacteria’, ‘Bacteroidetes’, ‘Actinobacteria’, especially OTUs from the same phyla (e.g. Proteobacteria) tended to co-occur more.

IV. CONCLUSIONS

Mining the marine microbial interaction patterns and diversity is a key for exploiting the marine resources. Considering the marine microbes are symbiosis or competition, exhibiting a numerous, significant intra- or inters- lineage associations, we used the network approach to analyze the potential interaction patterns among the marine microbes from the 16S rRNA sequences. The results show that the four seasonal marine microbial interaction networks have characters of complex networks, and the marine microbial interaction patterns are related with the seasonal variability.

Although we cannot claim that we have a comprehensive view of interactions within marine microbial communities, our network analysis method is more feasible and interesting for exploring the unseen patterns emerged in the complex dataset, including non-random association, deterministic processes at different taxonomic levels and expected relationship between community members.

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