Salt-tolerance Function in Halotolerant Bacteria from Coastal Soil and Sediment Present near Saltern Field of Hainan Island, China

Running title: salt-tolerance function in halotolerant bacteria from coastal soils and sediments

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Abstract

Understanding the salt-tolerance function in microbial communities may help illustrate the influences of salt concentration and other environmental factors on the soil biodiversity. Here, we combined high-throughput sequencing of 16S rDNA and ITS to investigate the distribution and salt-tolerance function of microbial communities in coastal soils and sediments present near the Yinggehai saltern field of Hainan Island, China. The microbial communities in the soil and sediments of the land zone (YGHLS), the intertidal zone (YGHIS), and the inshore zone (YGHWS) were compared. PCoA of weighted and unweighted UniFrac distances showed the difference in soil microbial communities across the samples. ANOSIM analysis revealed that the three samples were separated from each other. Halomonas, Halobacillus, and Wallemia presented in the samples and accounted for 0.0335 ± 0.0586%, 0.0241 ± 0.0304%, and 0.0308 ± 0.0445% of the total microbial community. Relative abundance of Na⁺:H⁺ antiporter, Trk system potassium uptake protein, and Kdp operon response regulator in the samples exceed 0.02%, 0.09%, and 0.06% respectively. It revealed that the halotolerant bacteria of Yinggehai coastal soil and sediment mainly take the Trk system for salt-tolerance function.

Keywords Salt-tolerance · coastal soils and sediments · saltern
Introduction

Island ecosystem is a unique and fragile ecosystem on earth. They provide special habitat for a wide variety of terrestrial and aquatic organisms and play important ecological and economic roles. Due to the high salt environment and other special environment factors, it owns abundant microbial genetic resources (Somboonna et al. 2012, Ng and Tan 2018, Sottorff et al. 2019). Characteristics of microbial community that exists in the coast near the saltern of an island is an interesting theme in ecology.

To assess the effects of hypersaline environment on microbial composition, many microbial biogeography studies examined coastal soils and sediments present in saltern fields. The phyla Proteobacteria, Firmicutes, and Actinobacteria were isolated from coastal solar salterns and revealed they have the ability to synthesize bioactive molecules (Boyadzhieva et al. 2018). Tang et al. (2011) reported 47 bacterial isolates either in the phylum Firmicutes or in the class Gammaproteobacteria isolated from a hypersaline pond in Sichuan, China. Characterization of heterotrophic prokaryote subgroups in the Sfax coastal solar salterns showed that the Alphaproteobacteria class in the non-crystallizer pond samples and the Bacteroidetes phylum in the crystallizer pond samples (Trigui et al. 2011). The different microbial genera present in the saline sediments and soils belonged mostly to halophilic bacteria, suggesting that the coast near saltern may be a valuable system to investigate saline influence the microbial community.

Yinggehai saltern is located in the southwest of Hainan Island, China. It was built in 1958, a high-salinity environment in the coastal waters. The saltern area is warm and hot all year and has very distinct wet and dry season. Evaporation is much greater than rainfall, and this superior climate condition has become a suitable place for salt drying. This study aims to explore the roles of the microbial community in the coastal soil and sediment near Yinggehai saltern by using 16S and ITS ribosomal DNA sequencing. In addition, we focus on salt-tolerance function in the microbial community of the soil and sediment. It will unravel the geographical factors affect the halotolerant bacteria community.

Materials and Methods

Study Site and Sample Collection

Yinggehai saltern is located in Hainan Island, China (Fig. 1). Three successional areas were selected for analysis, including the land zone (YGHLS, surrounded by plants, 18.4948N 108.7158E), the intertidal zone (YGHIS, ebb and flow of the tide, 18.4931N 108.7156E), and the inshore zone (YGHWS, immersed by
seawater, 18.4898N 108.7153E). The samples were collected in July 2018. Quadruplicate 1 × 1 m² plots were separated by 10 m in each area. For each plot, 5 soil cores (5–10 cm diameter × 5–10 cm depth) were taken to make a composite sample.

**Determination of Physicochemical Characteristics**

Soil and sediment samples were air-dried and sieved through a 100-mesh sieve for determination of organic matter (OM). Electrical conductivity (EC), pH, total nitrogen, sodium (Na), sulfate (S\(^{2-}\)), nitrate (N\(^{−}\)) and ammonium (N\(^{+}\)) were sieved through a 10-mesh sieve. The physicochemical characteristics were tested according to the method described by Chen et al. (2020).

**Quantification of Microbial Abundance**

OMEGA Soil DNA Kit was used to extracted genomic DNA. Quantity and quality of DNA was measured by NanoDrop ND-1000 spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA) and agarose gel electrophoresis. Microbial abundance quantification was evaluated by real-time quantitative polymerase chain reaction (Xiong et al. 2017). The copy number of 16S rRNA and ITS genes was determined by five replicates of each sample with the primers 338F (5′-ACTCCTACGGAGGCAGCA-3′) and 519R (5′-ATTACCAGGCTGCTGG-3′) (Zhang et al. 2017) and the primers ITS5F (5′-GGAAGTAAAAGTCTGAAACAAGG-3′) and ITS1R (5′-GCTGCGTTCTTCATCGATGC-3′) respectively.

**PCR Amplification and Sequencing**

The 16S rDNA V3–V4 region was amplified using primer 338F (5′-ACTCCTACGGAGGCAGCA-3′) and primer 806R (5′-GGACTACHVGGGTWTCTAAT-3′) (Hong et al. 2020), and the ITS1 region was amplified using primer ITS5 (5′-GGAAGTAAAAGTCTGAAACAAGG-3′) and primer ITS2 (5′-GCTGCGTTCTTCATCGATGC-3′) (White et al. 1990). The primers incorporated with sample specific 7 bp barcodes were used to perform multiplex sequencing. PCR mixtures contained 5 μl of Q5 reaction buffer (5×), 5 μl of Q5 High-Fidelity GC buffer (5×), 2 μl of template, 1 μl (10 μM) of each primer, 2 μl (2.5 mM) of dNTPs, 0.25 μl of Q5 High-Fidelity DNA Polymerase (5 U/μl), and 8.75 μl of ddH₂O. Reactions were programmed for a denaturation at 98 °C for 2 min, followed by 25 cycles consisting of 98 °C for 15 s, 55 °C for 30 s, and 72 °C for 30 s, with an extension of 5 min at 72 °C. PCR products were purified and quantified with Agencourt AMPure Beads (Beckman Coulter, Indianapolis, IN) and the PicoGreen dsDNA Assay Kit
(Invitrogen, Carlsbad, CA, USA) separately. Products were pooled in equal amounts, and sequencing was performed using the Illumina MiSeq platform at Shanghai Personal Biotechnology Co., Ltd.

**Sequence Analysis**

QIIME v1.8.0 was used for the sequencing data process (Caporaso et al. 2010). Valid sequences were identified based on raw sequencing reads with exact matches to the barcodes. Paired-end reads assembled using FLASH (Magoc and Salzberg, 2011). The high-quality sequences were assigned to operational taxonomic units (OTUs) by UCLUST with a 97% similarity threshold (Edgar 2010). Representative microbial sequences were selected from each OTU using default parameters, and matched against the SILVA and UNITE databases, respectively. It discarded the OTUs containing less than 0.001% of total sequences across all samples. An averaged, rounded rarefied OTU table was generated by averaging 100 evenly resampled OTU subsets under 90% of the minimum sequencing depth to minimize the difference in sequencing depth across samples.

**Bioinformatics and Statistical Analysis**

Chao1 richness estimator, Shannon diversity index, Simpson index, and ACE metric (abundance-based coverage estimator) were estimated using the OTU table in QIIME. Beta diversity analysis was performed using UniFrac distance metrics (Lozupone and Knight, 2005; Lozupone et al. 2006) and visualized via principal coordinate analysis (PCoA). Significant differences in microbial structures were evaluated based on analysis of similarity (ANOSIM) performed by the R package “vegan” (v3.2.0) (Clarke 1993). Venn diagrams were generated using the R package “VennDiagram” (Zaura et al. 2009). The linear regression analysis was conducted with Origin v 9.1. PICRUSt2 analysis was performed to gain insight for the salt-tolerance function (Langille et al. 2013). The gene family counts for each sample were derived from the KEGG ortholog (KO).

**Sequence Accession Numbers**

The raw sequence data are available in the NCBI Sequence Read Archive (SRA) database under the accession numbers PRJNA699381 and PRJNA699383.

**Results**

**Site Characteristics**

The selected sampling sites were from the land zone to the inshore zone, which is nearby Yinggehai saltern.
The variations of a wide range of physicochemical factors were observed. This analysis could reflect the ecosystem nearby the saltern zone (Table S1). The $\text{NH}_4^+$-N of the samples from the three sites did not vary much. The YGHWS sample had significant (Duncan test, $P < 0.05$) higher sodium ($2338.47 \pm 78.58$ mg/kg) and sulfate ($187.35 \pm 8.50$ mg/kg) contents. Detailed descriptions of the data are provided in Supplementary Table S1.

**Microbial Community Diversity and Structure**

The microbial communities could be adequately represented by the observed OTUs based on the relatively stable rarefaction curves (Fig. S1). Statistical results demonstrated that the most diversity indices were significantly different among the three sampling points at bacterial and fungal communities (Table 1). Although Shannon, evenness, and simpson index of bacterial community were not significantly different among the three sampling points, the other two diversity indices were significantly related to each other. Venn diagrams showed that the unique bacterial OUTs in the YGHLS, YGHIS, and YGHWS were 2118, 1587 and 2416, the unique fungal OUTs in the YGHLS, YGHIS, and YGHWS were 775, 716 and 1351 (Fig. S2). The difference in soil microbial communities across the samples was showed by PCoA (Fig. 2). ANOSIM analysis (Bray–Curtis) revealed that the three samples were separated from each other (Table S2, YGHWS vs YGHIS vs YGHLS, $P < 0.05$). In bacteria, YGHWS samples were not separated from group of the other two samples based on Bray–Curtis analysis (ANOSIM, YGHWS vs YGHIS and YGHLS, $P > 0.05$). Similarly, the fungi of YGHIS samples clustered tightly together with YGHLS and YGHWS group (ANOSIM, YGHIS vs YGHWS and YGHLS, $P > 0.05$).

**Microbial Community Composition**

The taxonomy of the bacterial community at the phylum and genus levels is presented in Fig. 3A and Fig. 3B. Proteobacteria, Actinobacteria, Acidobacteria, Chloroflexi, and Gemmatimonadetes were the five most abundant phyla in all samples in decreasing order and accounted for 84.41 ± 3.23% of the total microbial community. *Halomonas* and *Halobacillus* presented in the samples and accounted for 0.0335 ± 0.0586% and 0.0241 ± 0.0304% of the total microbial community, respectively. The identified phyla and genera of samples from the three sites were shown in Fig. 3A and Fig. 3B. Furthermore, the samples from three sites clustered independently in the bacterial genus distribution (Fig. S 3A).

Furthermore, we analyzed the relative abundance of fungi in the three samples at the phylum and genus
levels. The fungal phyla Ascomycota, Basidiomycota, Ciliophora, Zygomycota, Cercozoa, and Chytridiomycota were more abundant (Fig. 3C). Schizophyllum, Bullera, Cryptococcus, Talaromyces, Aspergillus, Tomentella, Spiromastix, and Auricularia were detected in all the samples (Fig. 3D). A rare genus Wallemia presented in the samples and accounted for 0.0308 ± 0.0445% of the total microbial community. The patterns of fungal genus distribution are showed in Fig. S3B.

**Relationship between Physicochemical Characteristics and Abundance of Halophilic Bacteria and Fungi**

Sodium and sulphate were selected from our initial physicochemical indicators in the linear model, showing that they did not have linear relationships with Halobacillus and Wallemia (Fig. 4). However, a bacterial genus Halomonas showed positive relationship with sodium ($R^2 = 0.501, P < 0.05$) and sulphate ($R^2 = 0.494, P < 0.05$) content. It indicated that the sodium and sulphate content significantly affected the Halomonas relative abundance, whereas they did not relate to Halobacillus and Wallemia.

**Relationship between Microbial Indicators and Halophilic Bacteria Abundance**

The structure (weighted PCoA1) and richness were selected as microbial indicators in the linear model, as they could explain the relationship with halophilic bacteria abundance. As a result, bacterial weighted PCoA1 ($R^2 = 0.065, P > 0.05$) and richness ($R^2 = 0.118, P > 0.05$) did not show linear relationships with Halobacillus and Halomonas abundance respectively. However, we found that bacterial weighted PCoA1 ($R^2 = 0.393, P < 0.05$) had a significant positive relationship with Halomonas abundance. Similarly, we also found that bacterial richness ($R^2 = 0.348, P < 0.05$) had a significant positive relationship with Halobacillus abundance (Fig. 5).

**Bacterial function prediction**

The salt-tolerance functional profile of composting microbiota was predicted by PICRUSt depending on KEGG database. Relative abundance of Na⁺:H⁺ antiporter, Trk system potassium uptake protein and Kdp operon response regulator in the samples exceed 0.02%, 0.09%, and 0.06% respectively (Fig. 6). Specifically, Na⁺:H⁺ antiporter ranged from 0.000398 ± 0.00009% to 0.0235 ± 0.00228% at YGHLs samples, from 0.00162 ± 0.000725% to 0.02394 ± 0.00209% at YGHIS samples, and from 0.00129 ± 0.00154% to 0.01726 ± 0.00237% at YGHWs samples. Other salt-tolerance Trk system potassium uptake protein ranged from
0.03924 ± 0.00158% to 0.07783 ± 0.00369% at YGHLS samples, from 0.04028 ± 0.00442% to 0.07283 ± 
0.00524% at YGHIS samples, and from 0.04772 ± 0.013% to 0.04958 ± 0.0023% at YGHWS samples.
Furthermore, Kdp operon response regulator ranged from 0.05131 ± 0.000778% to 0.06426 ± 0.00501% at 
YGHLS samples, from 0.04603 ± 0.00422% to 0.05506 ± 0.00962% at YGHIS samples, and from 0.03306 ± 
0.00228% to 0.03487 ± 0.0033% at YGHWS samples.

**Discussion**

Proteobacteria, Bacteroidetes, Chloroflexi, and Firmicutes dominated the bacterial communities of sediment 
samples present near the Mediterranean Sea coast of Camargue, while members belonging to Proteobacteria, 
Bacteroidetes, Gemmatimonadetes, Actinobacteria, Firmicutes, and Acidobacteria dominated the bacterial 
communities of the soil samples (Osman et al. 2018). In this study, Proteobacteria, Actinobacteria, 
Acidobacteria, Chloroflexi and Gemmatimonadetes were the five most abundant phyla in the samples.
Pomorie salterns and Burgas salterns located at Burgas Bay, Black Sea coast, Bulgaria. And, bacterial genera 
*Halomonas* was isolated from the two salterns (Boyadzhieva et al. 2018). Other genera *Salimicrobium*, 
*Halalkalibacillus*, *Virgibacillus*, *Alkalibacillus*, *Marinococcus*, *Halobacillus*, *Halomonas*, *Idiomarina*, 
*Chromohalobacter*, and *Halovibrio* were obtained from a hypersaline pond in Sichuan, China (Tang et al. 
2011). Fungal genera *Wallemia ichthyophaga* have been isolated globally from natural hypersaline 
environments (Zalar et al. 2005; Gunde-Cimerman et al. 2009). *Halomonas*, *Halobacillus* and *Wallemia* were 
also found in this study. It indicated that bacterial and fungal community composition located in Yinggehai 
coastal soil and sediment were similar with other hypersaline environments.

The observed shifts in salt tolerance along the gradients indicate that salinity environmental factors had 
shaped trait distributions and filtered the microbial communities. Previous studies revealed that 
physicochemical factors, such as, pH, salinity, exchangeable cations, C/N ratio, OM content, and sulfate 
content, they were major drivers that affected microbial community structure and function in the soil and 
sediment samples (Jorgensen et al. 2012; Dini-Andreote et al. 2014; Zeglin et al. 2016; Verma et al. 2017; 
Kang et al. 2018; Canini et al. 2020). Sodium and sulfate in the YGHWS sample showed significantly higher 
than the samples from other two site. Furthermore, the samples from three sites separated from each other in 
the weighted and unweighted UniFrac distances. Typically, content of sodium and sulphate showed a 
significantly positive relationship with halophilic bacteria *Halomonas* abundance. It revealed that the
halophilic bacteria abundance of coastal soils and sediments present near saltern field was possibly influenced by the soil physicochemical characteristics. However, *Halobacillus* and *Wallemia* did not show linear relationships with sodium and sulphate. The influence was obviously limited to the specific genus *Halomonas*. Interestingly, the bacterial weighted PCoA1 had a significant positive relationship with *Halomonas* abundance. Thus, *Halomonas* abundance was directly influenced by physicochemical characteristics or bacterial weighted PCoA1? It still needs more studies to ascertain.

The Na⁺/H⁺ antiporter is ubiquitous from bacteria to mammals and plays an important role in extrusion of Na⁺ or Li⁺ in exchange for H⁺ to keep the cytoplasm isosmotic with the environment and to avoid intoxication of living cells (Yang et al. 2006). At a high external salt concentration, bacteria take up K⁺ from the surrounding medium and accumulate K⁺ as an osmoregulatory solute to achieve an osmotic equilibrium. The major transport systems for K⁺ accumulation in these organisms is the transporter Kdp, the Ktr system, and the Trk transporter (Kraegeloh et al. 2005). Relative abundance of Trk transporter was higher than Na⁺:H⁺ antiporter and Kdp transporter in the samples. It indicated that the halotolerant bacteria of Yinggehai coastal soil and sediment mainly take the Trk system in response to Na⁺ osmotic stress.

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**Compliance with ethical standards**

**Conflict of Interest** The authors declare that they have no conflict of interest.

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