Metagenomics Reveals Dominant Unusual Sulfur Oxidizers Inhabiting Active Hydrothermal Chimneys From the Southwest Indian Ridge

Yong Wang1*, Hong-Yu Bi2†, Hua-Guan Chen2,3, Peng-Fei Zheng2, Ying-Li Zhou2,3 and Jiang-Tao Li4

1 Institute for Marine Engineering, Shenzhen International Graduate School, Tsinghua University, Shenzhen, China, 2 Institute of Deep Sea Science and Engineering, Chinese Academy of Sciences, Sanya, China, 3 College of Marine Sciences, University of Chinese Academy of Sciences, Beijing, China, 4 State Key Laboratory of Marine Geology, Tongji University, Shanghai, China

The deep-sea hydrothermal vents (DSHVs) in the Southwest Indian Ridge (SWIR) are formed by specific geological settings. However, the community structure and ecological function of the microbial inhabitants on the sulfide chimneys of active hydrothermal vents remain largely unknown. In this study, our analyses of 16S rRNA gene amplicons and 16S rRNA metagenomic reads showed the dominance of sulfur-oxidizing Ectothiorhodospiraceae, Thiomicrorhabdus, Sulfurimonas, and Sulfurovum on the wall of two active hydrothermal chimneys. Compared with the inactive hydrothermal sediments of SWIR, the active hydrothermal chimneys lacked sulfur-reducing bacteria. The metabolic potentials of the retrieved 82 metagenome-assembled genomes (MAGs) suggest that sulfur oxidation might be conducted by Thiohalomonadales (classified as Ectothiorhodospiraceae based on 16S rRNA gene amplicons), Sulfurovaceae, Hyphomicrobiaceae, Thiotrichaceae, Thiomicrospiraceae, and Rhodobacteraceae. For CO2 fixation, the Calvin-Benson-Bassham and reductive TCA pathways were employed by these bacteria. In Thiohalomonadales MAGs, we revealed putative phytochrome, carotenoid precursor, and squalene synthesis pathways, indicating a possible capacity of Thiohalomonadales in adaptation to dynamics redox conditions and the utilization of red light from the hot hydrothermal chimneys for photolithotrophic growth. This study, therefore, reveals unique microbiomes and their genomic features in the active hydrothermal chimneys of SWIR, which casts light on ecosystem establishment and development in hydrothermal fields and the deep biosphere.

Keywords: deep-sea hydrothermal vents, SWIR, thiohalomonadales, metagenomics, red light photosynthesis

INTRODUCTION

Deep-sea hydrothermal vents (DSHVs) are located in tectonically active areas where plate boundaries move at different speeds along mid-ocean ridges. DSHV is an important conduit for the exchange of energy and materials between the Earth’s interior and the ocean. Since the first report in 1977, DSHVs as a deep-sea extreme environment have attracted great concerns about the microbial extremophilic inhabitants with respect to early life form, chemosynthesis, and adaptation (Corliss et al., 1979; Francheteau et al., 1979; Jannasch and Mottl, 1985;
Kelley et al., 2005; Dick, 2019). The mixture of cold, oxic deep-sea water, and highly reducing fluids with high concentrations of hydrogen, sulfide, and methane was ideal for chemosautrophs and CI oxidizers (Dick et al., 2013). The complex dynamic habitats have steep thermal and chemical gradients, and different microbial populations are simultaneously involved in many important biogeochemical processes, such as the nitrogen, sulfur, and carbon cycles that also occur in symbionts of megafauna around DSHV (Jannasch and Mottl, 1985; Connelly et al., 2012). Strong dynamics of geochemistry and temperature provide a wide range of habitats for deep-sea microorganisms, and therefore, niche specificity of microbes has been demonstrated in different environmental settings by previous studies (Dick, 2019 and references therein).

Sulfide samples from inactive and active hydrothermal vents are distinct in microbial community structure and ecological function in the Indian Ocean (Zhang et al., 2016; Han et al., 2018; Adam et al., 2020; Hou et al., 2020). Sulfur oxidizing bacteria (SOB) dominate various hydrothermal sediments and flumes and are classified to be aerobes (e.g., SUP05 and Beggiatoa from Gammaproteobacteria), microaerobes (e.g., Sulfurimonas and Sulfurovum from Epsilonbacteraeota), and anaerobes (e.g., Caminibacter and Nautila from Epsilonbacteraeta) (López-Garcia et al., 2003; Dick, 2019; Meier et al., 2019; Hou et al., 2020) with preference to different electron donors. For the microaerobic SOB, the cbb3-type cytochrome c oxidase was involved in adaptation to low oxygen concentrations for respiration (Hou et al., 2020; Dong et al., 2021); however, the mechanism for reducing the damage by high oxygen flux is largely unknown. Considering the highly variable microenvironments adjacent to an active hydrothermal vent, there are perhaps much more SOB species that have evolved to adapt to the varying temperature and redox conditions. A photoautotrophic bacterium has been isolated from a hydrothermal chimney (Beatty et al., 2005). Evidence shows that this bacterial isolate from the Prosthecocloris genus can absorb weak ultra-red light as energy to fix CO$_2$ and oxidize H$_2$S. The abundance and distribution of such photoautotrophic SOB in the dark ocean are still unclear up to date.

Since the first report in the Southwest Indian Ridge (SWIR) more than a decade ago (German et al., 1998), multiple hydrothermal fields have been discovered in the Indian Ocean (Van Dover et al., 2001). Low-temperature hydrothermal activity because of the effect of an ultra-slow spreading speed of tectonic plates was reported in the SWIR (Tao et al., 2012). The discovery of hydrothermal fields in such an ultra-slow spreading ridge provides an unprecedented opportunity to understand microbially mediated biogeochemistry. A serpentinite-hosted hydrothermal site and distinct microbial community resembling those of the Lost City hydrothermal field in the Atlantic Ocean were reported in a magma-poor area of the eastern SWIR (Lecoeuvre et al., 2021). The relatively low temperature and serpentinitization of SWIR hydrothermal fields (Zhou and Dick, 2013) had probably shaped the microbial inhabitants in the hydrothermal fields. However, microbial genomics studies on the SWIR hydrothermal areas are still limited. *Thiomicrothrix* has been reported as a dominant SOB in SWIR (Cao et al., 2014), and was also isolated from the other hydrothermal vents (Brazelton and Baross, 2010; Scott et al., 2019). A recent study revealed a large number of Gammaproteobacteria with sulfur oxidation potentials using nitrate and oxygen as electron acceptors in the SWIR inactive hydrothermal sediments (Dong et al., 2021). We employed metagenomics to compare the microorganisms in the active hydrothermal vents with those from inactive ones to reveal microbial community structure and ecological function driven by active “black-smoker” hydrothermal seepage in SWIR. We hereby report a unique microbiome exclusively composed of Thiohalomonadales-dominated SOB, which has not been reported in DSHVs and presumably plays a critical role in the initiation of the deep-sea hydrothermal ecosystem in SWIR.

**MATERIALS AND METHODS**

**Sample Collection and Mineral Analysis**

During the *R/V* DY35 cruise, the manned submersible “jiaolong” collected sulfide samples from hydrothermal chimneys by dives no. 96 (49.65°E, 37.78°S; depth: 2,768) and no. 100 (49.65°E, 37.78°S; depth: 2,755 m) in SWIR (Figure 1, Supplementary Table I, and Supplementary Figure 1). A sulfide chimney sample was obtained from an inactive hydrothermal vent (49.63°E, 37.77°S; depth: 2,789 m) by dive no. 95 at a depth of 2,789 m. The samples were stored in sterile bags and maintained at −80°C until use. The temperature and pH of the hydrothermal fluid were measured *in situ* with a Miniature Autonomous Plume Recorder (MAPR) (Baker and Milburn, 1997).

An X-ray diffraction (XRD) analysis was carried out to determine the major minerals of the sulfide chimney using an Empyrean X-ray diffractometer (PANalytical, Malvern, United Kingdom). The detection parameters were Cu Kα radiation at 45 kV and 40 mA; Goniometer PW3050/60 (Theta/Theta); scanning from 3° to 85° with 0.03 step size (°2Th). The mineral components were converted to wt%.

**High-Throughput Sequencing and Analyses of 16S rRNA Gene Amplicons**

Genomic DNA of sulfide samples was extracted from a 2 g sample using the MO BIO Powersoil DNA isolation kit (Qiagen, Carlsbad, CA, United States). The quality and quantity of the DNA extractions were checked by the Qubit 2.0 fluorometer (Invitrogen, Carlsbad, CA, United States). The quality and quantity of the sample using the MO BIO Powersoil DNA isolation kit (Qiagen, Hilden, Germany) was measured separately, and were sequenced on an Illumina Miseq sequencer.

Analyses of 16S rRNA Gene Amplicons

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FIGURE 1 | Sampling site at “Longqi” hydrothermal field. During RV DY35, two chimney rocks (D96S and D100S) were obtained by the “Jiaolong” submersible from active sulfide chimney bodies (inset figures) at “Longqi” hydrothermal field in the Southwest Indian Ocean. A sulfide sample (D95A) was also collected from a nearby inactive chimney by the submersible. The details of the samples are listed in Supplementary Table 1.

to obtain $2 \times 300$ bp paired-end sequencing reads (Illumina, San Diego, CA, United States).

The sequencing reads of 16S rRNA gene amplicons were checked by FASTQC (v0.11.8)$^1$ and merged by the QIIME 1 workflow (Kuczynski et al., 2011). The adaptors and low-quality reads were trimmed by the NGS QC Toolkit (v2.3.3) (Patel and Jain, 2012). Operational taxonomic units (OTUs) at 97% similarity were selected between the qualified reads by the QIIME 1 workflow (Kuczynski et al., 2011). Taxonomic classification of the OTUs was performed with vsearch against the SILVA database v138 (Quast et al., 2013). A principal component analysis (PCoA) was performed to estimate the similarity between the microbial communities at the family level using vegan in the R package (Dixon, 2003).

Metagenomics Analyses of Microbial Community and Metabolic Potentials

Raw reads of metagenomes were qualified by removing adapters and then filtered by using fastp (version 0.20.0) (Chen et al., 2018) with parameters (-w 16 -q 20 -u 20 -g -c -W 5 -3 -l 50). Low-quality reads (assigned by a quality score $< 20$ for $>20\%$ of the read length), and those shorter than 50 bp and unpaired were removed. Metagenomics reads that were mapped onto sequences in an in-house contaminant database (such as, sequences of human, mouse, and common laboratory contaminant bacterial genomes downloaded from the National Center for Biotechnology Information [NCBI]) (Huang et al., 2019) by Bowtie2 (version 2.4.1) (Langmead and Salzberg, 2012) were discarded.

Furthermore, 16S miTags were extracted from qualified metagenomic reads using rna_hmm3.py (Huang et al., 2019), which employs HMMER (version 3.1b2) to predict ribosomal RNA gene fragments from both forward and reverse metagenomic reads. An in-house python script was used to extract 16S miTags. The 16S miTags were imported into Qiime1 with a setting of –type “SampleData[Sequences]” and dereplicated redundancy to yield representative 16S miTag sequences. The classify-consensus-vsearch command in QIIME v1.9.1 (Kuczynski et al., 2011) was used to classify the representative 16S miTags using the SILVA database.
SSU database v138 as a reference (Quast et al., 2013), as mentioned above.

The qualified reads were assembled using SPAdes (v3.13) (Nurk et al., 2013) with a kmer set of 21, 33, 55, 77, 99, and 127 under the “—careful” mode (metagenome mode). Genome binning from assembled contigs > 2 kbp was performed by running three tools MaxBin (Wu et al., 2016), MetaBAT (Kang et al., 2019), and CONCOCT (Alneberg et al., 2014) using their default settings. Raw genome bins resulting from the three approaches were combined, followed by a selection of the best genome for each genome set using the bin_refinement module in metaWRAP (v1.2) (Uritskiy et al., 2018). During the bin refinement, we applied CheckM_lineage (v1.0.12) (Parks et al., 2015) to evaluate completeness and contamination for each bin. The draft genomes with >50% completeness and <10% contamination were retained for further analysis. Metagenome-assembled genomes (MAGs) were treated with dRep software (Olm et al., 2017) to dereplicate the MAGs by an average nucleotide identity (ANI) threshold of 95% (dereplicate -p 40 -comp 50 -con 10 -pa 0.95 -sa 0.95 -l 10000 -nc 0.30). The taxonomic position of the genomes was identified using genome taxonomy database (GTDB)-tk v0.2.2 (Chaumeil et al., 2020), as well as the calculation of relative evolutionary distance (RED).

Coding regions (CDS) for individual MAGs were predicted using Prodigal (version v2.6.3) (Hyatt et al., 2010) with option “-p meta.” Annotation of CDSs was performed using KofamScan (version 1.1.0) (Aramaki et al., 2019) and using Blastp (E value = 1e-5) against the Clusters of Orthologous Gene (COG) and NCBI_nr databases. The results were visualized using the heatmap package of the R platform.

Phylogenetic Tree Construction

For phylogenetic analysis of Thiohalomonadales genomes, 43 conserved proteins were obtained by the CheckM program with default settings (Katoh et al., 2009) and were used for alignment with Mafft (v7.g53, setting: –maxiterate 1000-localpair) (Katoh and Standley, 2013), followed by a further optimization with trimAl (v1.4) (Capella-Gutiérrez et al., 2009). A maximum likelihood (ML) tree was reconstructed using IQ-TREE (v1.6.12) (Parks et al., 2015) with the “LG + F + R6” model (Price et al., 2017) with a kmer set of 21, 33, 55, 77, 99, and 127 under the “—careful” mode (metagenome mode). Genome binning from assembled contigs > 2 kbp was performed by running three tools MaxBin (Wu et al., 2016), MetaBAT (Kang et al., 2019), and CONCOCT (Alneberg et al., 2014) using their default settings. Raw genome bins resulting from the three approaches were combined, followed by a selection of the best genome for each genome set using the bin_refinement module in metaWRAP (v1.2) (Uritskiy et al., 2018). During the bin refinement, we applied CheckM_lineage (v1.0.12) (Parks et al., 2015) to evaluate completeness and contamination for each bin. The draft genomes with >50% completeness and <10% contamination were retained for further analysis. Metagenome-assembled genomes (MAGs) were treated with dRep software (Olm et al., 2017) to dereplicate the MAGs by an average nucleotide identity (ANI) threshold of 95% (dereplicate -p 40 -comp 50 -con 10 -pa 0.95 -sa 0.95 -l 10000 -nc 0.30). The taxonomic position of the genomes was identified using genome taxonomy database (GTDB)-tk v0.2.2 (Chaumeil et al., 2020), as well as the calculation of relative evolutionary distance (RED).

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RESULTS

Mineral Composition of the Southwest Indian Ridge Hydrothermal Chimneys

The “Longqi” hydrothermal area is located in the Southwest Indian Ridge. Two sulfide samples, D96S and D100S, were collected from the outer wall of active hydrothermal chimneys on 14 January and 4 February 2015 by the “Jiaolong” submersible (Figure 1 and Supplementary Table 1) in dives nos. 96 and 100, respectively. The chimney rocks were about 10 cm away from the vents. Strong seepage from the hydrothermal vent could be observed. The temperature of the hydrothermal fluid was recorded to be 362 and 365°C for the vents near D96S and D100S, respectively, while the bottom seawater was ∼2.2°C. The pH of the fluid from the two vents was between 3.47 and 3.58. The surface of the D96S sample is gray and has been partially oxidized to grayish brown, while it was light yellow inward with metallic luster (Supplementary Figure 1). There was no megafauna attached to these chimneys. An XRD analysis revealed that D95S and D100S were exclusively composed of sphalerite (ZnS) with a small amount of pyrite (FeS₂) (<1%) (Supplementary Figure 2).

Oxidized minerals were not present in the XRD results, which is in contrast to the dominance of copper-containing chalcopyrite (CuFeS₂) and kusaschiite (CuBi₂O₄) in inactive hydrothermal sediments of SWIR (Cao et al., 2014).

Analysis of Prokaryotic Community Structure

Prokaryotic communities in our chimney samples were examined by sequencing and analysis of 16S rRNA gene amplicons and 16S rRNA gene tags in metagenomic reads (16S miTags), and were then compared with those of the sulfide samples from other SWIR sites (Supplementary Table 1; Dong et al., 2021). There are a total of 30,189 qualified 16S rRNA gene amplicons for the characterization of microbial communities. They were then clustered with 97% similarity into 6,994 OTUs (Supplementary Table 2). Proteobacteria dominated all the SWIR sulfide samples of this study (Figure 2A), except for D95S, in which Bacteroidetes (20%) were the most prevalent phylum. Epsilonbacteraeota was particularly enriched in the chimney samples, and Archaea was not present in these communities. At the family level, Ectothiorhodospiraceae (average 11.9%) and Thiovulaceae (average 6.8%) were the most abundant families in both the 16S rRNA gene amplicon- and miTag-based communities of the D96S sample. Ectothiorhodospiraceae is significantly more abundant in the active chimney samples (D96S and D100S) than in other SWIR hydrothermal sediments using the U-test (p < 10⁻¹⁰). In contrast, D100S contains a high proportion (7.7–19.1%) of Sulfurovaceae, which outcompetes Ectothiorhodospiraceae (~6.2%) and Thiovulaceae (~3.2%) as the dominant SOB (Figure 2B). Sulfur reducing bacteria (SRB) represented by Desulfobulbaceae and Thermodesulfovibrionaceae are abundantly present only in D95S and the reference SWIR sulfide samples. Thermodesulfovibrionaceae (Nitrospirae) occupied 6.3% of the community in D95S. The distinct distribution of Ectothiorhodospiraceae and Desulfobulbaceae between active chimney samples (D96S and D100S) and other hydrothermal samples from the SWIR was further illustrated by a chord diagram (Figure 3). The plotting further revealed special enrichment of Thiotrichaceae as the major SOB in inactive hydrothermal sediments, particularly in the S12T1 sample. A PCoA plot separated the microbial communities of D96S and D100S from the references and D95S. For D96S and D100S, the community structures revealed by the 16S rRNA gene amplicon and miTags were consistent with respect to the PCoA clustering (Figure 4).

Metagenomics Analysis of Dominant Species Inhabiting Active Hydrothermal Chimney

To understand the ecological role of microbial inhabitants in the active hydrothermal vents, we obtained 12.1 Gb raw
FIGURE 2 | Microbial community structures of hydrothermal samples. Taxonomic classification of operational taxonomic units (OTUs) at 97% similarity was performed by comparing with reference sequences of SILVA database v138. The microbial community structures were shown at phylum (A) and family (B) levels. D100SA, D96SA1, D96SA2, and D95SA refer to 16S rRNA gene amplicons; D100SM and D96SM are 16S miTags. The other samples are referred to Supplementary Table 2.
Illumina Miseq paired-end sequencing data (2 × 300 bp) and retained 8.3 Gb clean data for subsequent metagenomics work of D96S and D100S (Supplementary Table 3). After assembly and genome binning, a total of 82 MAGs were retrieved from D96S and D100S metagenomic assemblies (Supplementary Table 4), among which 57 MAGs were of good quality (completeness > 80% and contamination < 5%). Taxonomic classification of the MAGs against GTDB indicates that they were affiliated with 12 prokaryotic phyla mainly composed of Proteobacteria (n = 39), Bacteroidota (n = 20), Campylobacterota (n = 7), and Desulfobacterota (n = 5; Supplementary Table 4).

The abundant *Ectothiorhodospiraceae* in D96S and D100S identified by 16S rRNA gene amplicon sequencing were not present in the taxa of the MAGs. By searching the 16S rRNA gene amplicons of *Ectothiorhodospiraceae* against those extracted from the MAGs (>97% similarity), we reassigned *Ectothiorhodospiraceae* to the orders Thiohalomonadasaes SZUA-152 (n = 8) and SZUA-140 (n = 1) of Gammaproteobacteria in GTDB taxonomy. In a phylogenomics tree, these MAGs were clustered with the genomes obtained from hypersaline soda lake sediment, ground water, and other deep-sea hydrothermal vents, such as Mid-Atlantic Ridge, East Pacific Rise, and Lau Basin.
of Tonga (Figure 5). Three phylogenetic groups were formed with most of our MAGs in Thiohalomonadales SZUA-152, a new order in GTDB.

On the basis of Kyoto Encyclopedia of Genes and Genomes (KEGG) gene annotation of the 57 good-quality MAGs, the functional genes for carbon, nitrogen, and sulfur metabolisms mediated by the major families of the chimney inhabitants were examined to predict their ecological functions. Hydrothermal vents discharge large amounts of hydrogen, methane, and hydrogen sulfide to deep waters (Jannasch and Mottl, 1985; Kawagucci et al., 2010; Dick, 2019). Thiosulfate oxidizing genes soxABXYZ, and sulfide oxidizing genes dsrAB, fccAB, and sqr were found in ≥50% of the MAGs for Thiohalomonadales (Gammaproteobacteria), Thiomicrospiraceae (Gammaproteobacteria), and Rhodobacteraceae (Alphaproteobacteria) MAGs from D96S; and in Thiohalomonadales, Sulfurovaceae (Campylobacterota), Hyphomicrobiaceae (Alphaproteobacteria), and Thiotrichaceae (Gammaproteobacteria) MAGs from D100S (Figure 6). In the periplasmic space, FccAB catalyzes oxidation of sulfide to elemental sulfur with electrons being transferred to cytochrome c; the SQR enzyme oxidizes sulfide to polysulfide (Schutz et al., 1999). The DsrAB encoded by the SOB carry out sulfide oxidation to sulfite (Anantharaman et al., 2014; Dong et al., 2021). This indicates the capacity of sulfide oxidation to $S^0$, polysulfide or sulfate by these SOB bacteria inhabiting our active chimney samples under different fluxes of electron acceptors. The Rhodobacteraceae MAGs also harbor nitrate reduction genes narG, suggesting the coupling of nitrate reduction and sulfide oxidation. Sulfurimonadaceae and Sulfurovaceae encoded the reverse tricarboxylic acid (rTCA) pathway for autotrophic CO$_2$ fixation. The other SOB, with the exception of Thiotrichaceae, may rely on the Calvin–Benson–Bassham (CBB) pathway for autotrophs, as ribulose-1,5-bisphosphate carboxylase (RuBisCO) genes rbcLS were found in their MAGs. Wood–Ljungdahl pathway as an alternative autotrophic process was only encoded by Desulfobacteraceae MAGs. Acetate metabolism is highly required by the bacterial inhabitant, as evidenced by the prevalence of acetate assimilatory genes pta, acs, and ackA in these MAGs. Genomics data also indicate that Gammaproteobacteria might use hydrogen released from the vents as alternative energy sources for carbon fixation and metabolic activities as indicated by the detection of [NiFe]-hydrogenase genes.
Metabolism Potentials of Thiohalomonadales

We next examined the metabolic potentials and adaptive strategy of the dominant SOB, Thiohalomonadales, from our active hydrothermal vents using the annotation result of their five high-quality MAGs. The Thiohalomonadales MAGs harbor a complete set of sulfur-oxidizing genes, such as \textit{soxABXYZ}, \textit{fccABV}, \textit{sqr}, \textit{rdsrAB}, and \textit{TST}, that may yield $S_0$, polysulfide, or sulfate (Figure 7). Nitrate assimilatory reduction may occur in the Thiohalomonadales as the MAGs contain \textit{napAB} periplasmic...
nitrate reduction genes and nitrite reduction genes (nirBD) that are involved in ammonia production. Thiohalomonadales might also rely on the CBB pathway for CO₂ fixation. Thiohalomonadales MAGs contain cbb3 genes coding for phosphorylation respiration complex IV to overcome low oxygen conditions. Considering the high hydrogen content of the hydrothermal fluids, the microbes on the chimney are expected to be able to utilize hydrogen as an energy source. [NiFe] hydrogenase genes were present in the MAGs, along with hup genes responsible for the uptake and/or export of hydrogen. The Rnf complex as a cross-membrane proton pumping machine for the balance of cytoplasmic pH was encoded by the MAGs (Buckel and Thauer, 2013).

We identified all the related genes encoding the enzymes that participate in terpenoid backbone biosynthesis. Using geranylgeranyl diphtathate, Thiohalomonadales might generate phytoene, since the ctb gene encoding 15-cis-phytoene synthase was identified in the MAGs. The CrtB protein of Thiohalomonadales is most similar (82%) to a homolog of Thiorthicichaeae from a subseafloor aquifer. Phytoene is a precursor of zeta-carotene and can be catalyzed by phytoene desaturase Ctr1 for carotene production (Lang et al., 1994). However, ctr1 gene is missing from the MAGs, and the capacity of carotene biosynthesis by these SOB is thus questioned. The ctb gene was also present in the MAGs from RBG-16-57-12, Mariproducinae, Thiohalomonadales SZUA-140, and Melioribacteraceae. Most of these ctb-bearing MAGs also contain a farnesyl-diphosphate farnesyltransferase coding gene (FDFT1) that functions in squalene synthesis using farnesyl diphtathate.

In the Thiohalomonadales MAGs, a cph2 type bacteriophytochrome coding gene was identified to be 45% similar to a homolog from Pseudomonas aeruginosa. However, the putative bacteriophytochrome is featured with GGDEF and EAL domains but lacks GAF and PHY domains, indicating a discovery of a novel or malfunctional bacteriophytochrome (Gourinchas et al., 2019). The co-factor of bacteriophytochrome is biliverdin with a tetrapyrrolic structure. Heme synthesis pathway and biliverdin-producing heme oxygenase have been identified in the MAGs, which are vital for the potential function of bacteriophytochrome under red and far-red lights for Thiohalomonadales species (Vuillet et al., 2007; Gourinchas et al., 2019).

DISCUSSION

Discovery of Novel Microbiomes in Active Hydrothermal Chimenys

In this study, the “jiaolong” manned submersible with robotic precision was employed to locate the vent structure of sulfide chimneys in the active SWIR hydrothermal areas. Using these chimney samples, we report the genomes of dominant microbial species and the prevalence of distinct SOB affiliated with Campylobacterota, Gammaproteobacteria, and Alphaproteobacteria inhabiting the sulfide chimney, all of which have been reported in deep-sea cold seeps recently (Li et al., 2021). The microbial community in the active SWIR hydrothermal vents was remarkably different from those from the SWIR inactive hydrothermal sulfide sediments (Cao et al., 2014; Dong et al., 2021) and others DSHVs (Dick, 2019; Hou et al., 2020; Zeng et al., 2021). Culturable Thiohalomonadales SOB, such as Thiohalomonas denitrificans had been isolated from the hydrothermal vents of the Suiyo Seamount in the Pacific Ocean (Mori et al., 2015). In addition, this study demonstrates their distribution in the DSHVs located in the Mid-Atlantic Ridge, East Pacific Rise, and Lau Basin. Given the finding of their relatives in other worldwide sites, the dominance of this order in DSHVs was, however, not reported previously (Flores et al., 2011; Urich et al., 2014; Meier et al., 2017). Dominant SOB differed even between D96S and D100S, owing to different combinations of Thiohalomonadales, Thiomicrothrixbus, Sulfuravum, and Sulforimonas. This suggests a high diversity of SOB among active hydrothermal vents. Our genomics data predict other potential SOB, such as Hyphomicrobiaceae, Thiotrichaceae (represented by Beggiatoa), and Rhodobacteraceae in our samples, which have been rarely reported in other DSHVs. Considering the highly variable microenvironments adjacent to DSHVs, the composition of SOB inhabitants is predicted to be slightly diversified as reported for D96S and D100S by this study. In the late stages of hydrothermal vents, weak seepage of reducing fluids allows soaking of sulfide with oxic bottom water, which will gradually result in sulfide mineral oxidation and subsequent microbiob mediated reduction processes (Li et al., 2017). The microbiomes of D96S and D100S were solely constituted by SOB, which is an indicator of the initial stage of a hydrothermal ecosystem. This has not been noticed in the SWIR and even global DSHVs.

The strong hydrothermal venting probably creates a reducing environment that covers the chimney, prohibiting oxidation of the chimney sulfide in this study. The mineral components in our samples indicate an early stage of the chimney formation at D100S, as there were not any oxidized mineral components. As a result, we could not detect SRB in D100S, while in contrast, prevalent Thermodesulfovibrioaceae (SRB) was present in D95S. Although the sulfate concentration and nutrients of the samples were not analyzed due to their contact with sea water, we speculate that sulfate from sea water and produced by SOB might probably fuel the SRB in D95S. It seems that Sulfurovaceae, Thiohalomonadales, and Thiiovulaceae were distributed differently in the hydrothermal chimneys possibly due to selection of different SOB families by local environmental variants. For example, Sulfurovum is more tolerant to oxygen (Meier et al., 2017) and might prefer a more oxic chimney with lower impact of hydrothermal fluid. In this study, Thiohalomonadales probably employed cbb3 to cope with low oxygen; however, hypoxic exposure may also impair anaerobic and microaerobic microbial inhabitants (Lu and Imlay, 2021). We found that squalene as a bacterial hopanoid was likely synthesized by Thiohalomonadales, which is probably an efficient mechanism to scavenge single oxygen that may damage cell lipid by peroxidation (Kohno et al., 1995). The microbes in D96S and D100S have probably evolved to contain distinct gene profiles for adaptation to environmental changes due to the dynamics of gas and metal fluxes near active hydrothermal vents.
Photolithoautotrophic Potential of Sulfur-Oxidizing Thiohalomonadales

Ectothiorhodospiraceae was previously a family of Chromatiaceae, but was recently reclassified into a new family of Thiohalomonadales. Members of the Ectothiorhodospiraceae are versatile in metabolisms, including photolithotrophic, photoheterotrophic, chemoheterotrophic, chemolithotrophic, and methylotrophic bacteria using various electron acceptors, such as nitrite, sulfur compounds, and arsenite (Hallberg et al., 2011; Slobodkina et al., 2016). So far, all isolates from hydrothermal fields belonging to this family are sulfur-oxidizing bacteria (Imhoff, 2006; Mori et al., 2015; Slobodkina et al., 2016). In this study, we discovered three phylogenetic groups in Thiohalomonadales belonging to SZUA152, SZUA140, and an unclassified clade. Ectothiorhodospiraceae members are able to capture red/far-red light for anaerobic photosynthesis, by which sulfide is oxidized to elemental sulfur and deposited in periplasm for further oxidation to sulfate (Imhoff, 2006). Bacteriophytochromes are photosensitive proteins employed by bacteria to capture red/far-red light to mediate growth (Vuillet et al., 2007; Gourinchas et al., 2019). At hydrothermal vents with a temperature of \( \sim 350^\circ C \), most of the ambient light was detected to fall into near-infrared spectra between 750 and 1,050 nm (Dover et al., 1994). A recent report showed the promoted growth of a bacterial cultivation isolated from the Western Pacific DSHV under stimulation of infrared light at 940 nm (Liu et al., 2021). This is highly supported by a recent report that showed the presence of proteorhodopsin synthesis genes and photoautotrophic microbial groups in some metagenomes of active hydrothermal vents of SWIR (Chen et al., 2022). If the bacteriophytochromes and cofactor biliverdin are functional in Thiohalomonadales, they can probably sense red/near-infrared light from the hydrothermal vents to activate photosynthesis of Thiohalomonadales. In photosynthetic and non-photosynthetic organisms, carotenoids are synthesized to prevent the photooxidative damage (Glaeser and Klug, 2005). In the present study, although the gene coding for the enzyme catalyzing the final step of zeta-carotenoid was not found, there are still perhaps unknown genes responsible for carotenoid synthesis in the genomes. Aside from serving as light sensors, some bacteriophytochromes can be redox sensors to monitor the environmental changes, particularly in the environment approximate to hydrothermal vents where hot reductive fluid meets cold oxic sea water. Whether Thiohalomonadales can synthesize the \textit{bona fide} bacteriophytochromes for photoautotrophy warrants future experimental efforts using cultivated strains of this order from DSHVs.

Overall, we identified an unusual, uncultivated sulfur-oxidizing bacterial group that is particularly prevalent in active hydrothermal chimneys. Their potential capacity of sulfur oxidation and photoautotrophic lifestyle casts lights on the possible relationship with ancient lineages inhabiting the early earth with similar hydrothermal environments billions of years ago.
 ago (Lunine, 2006). The finding of this SOB group can probably provide insights into the future evolutionary study of ancient microbial lineages under the climate change in the long history of the Earth.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

**AUTHOR CONTRIBUTIONS**

H-YB and YW conceived the study and wrote the manuscript. H-GC, P-FZ, and Y-LZ performed the experiments. YW, H-YB, P-FZ, and Y-LZ analyzed the data and summarized the results. J-TL critically revised the manuscript. All authors contributed to the article and approved the submitted version.

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**FUNDING**

This study was supported by the Major Scientific and Technological Projects of Hainan Province (ZDKJ2021028).

**ACKNOWLEDGMENTS**

We thank the cruise members of D/V DY35 and “Jiaolong” submersible pilots for their efforts in sampling. We appreciate the Supercomputing Center of University of Sanya for providing the computation assistance.

**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2022.861795/full#supplementary-material
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