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<td>Author(s)</td>
<td>Li, H; Chen, S; Mu, BZ; Gu, JD</td>
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Molecular Detection of Anaerobic Ammonium-Oxidizing (Anammox) Bacteria in High-Temperature Petroleum Reservoirs

Hui Li • Shuo Chen • Bo-Zhong Mu • Ji-Dong Gu

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Abstract Anaerobic ammonium-oxidizing (anammox) process plays an important role in the nitrogen cycle of the worldwide anoxic and mesophilic habitats. Recently, the existence and activity of anammox bacteria have been detected in some thermophilic environments, but their existence in the geothermal subterranean oil reservoirs is still not reported. This study investigated the abundance, distribution and functional diversity of anammox bacteria in nine out of 17 samples. Most of 16S rRNA gene phylotypes are closely related to the known anammox bacterial genera Candidatus Brocadia, Candidatus Kuenenia, Candidatus Scalindua, and Candidatus Jettenia, while hzo gene phylotypes are closely related to the genera Candidatus Anammoxoglobus, Candidatus Kuenenia, Candidatus Scalindua, and Candidatus Jettenia. The total bacterial and anammox bacterial densities were 6.4±0.5×10^3 to 2.0±0.18×10^5 cells ml^-1 and 6.6±0.51×10^2 to 4.9±0.36×10^4 cell ml^-1, respectively. The cluster I of 16S rRNA gene sequences showed distant identity (<92%) to the known Candidatus Scalindua species, inferring this cluster of anammox bacteria to be a new species, and a tentative name Candidatus “Scalindua sinooilfield” was proposed. The results extended the existence of anammox bacteria to the high-temperature oil reservoirs.

Introduction

A petroleum reservoir is a typically extreme environment characterized by high temperature, high pressure, high salinity, and strictly anoxic condition. From the first description of the microorganisms in oil deposits [4], the microbiology of petroleum reservoirs subsequently developed along with petroleum exploration. A wide range of microorganisms have been cultured from oilfields with sulfate reduction, fermentation, acetogenesis, and methanogenesis as major metabolic types [40, 63]. The culture-independent researches indicated more diverse microbes than previously perceived in oilfields worldwide [1, 5, 7, 12, 15, 28, 33, 41, 42, 46]. Recently, much more attention has been paid to the microbial flora associated with the nitrogen cycle in the subterranean petroleum-rich strata because the source of nitrogen is a limiting factor to the microbial community activity under the oilfield condition. In the petroleum industry, substantial amounts of nitrate were injected into the oil reservoirs in an attempt to control the souring [21] through changing the microbial community structure in the subsurface strata. Several nitrogen-cycle-
related microorganisms were activated by the injected nitrate, including the nitrate-reducing independent with sulfite oxidation bacteria and heterotrophic nitrate-reducing bacteria [20]. Along with these processes, nitrite and ammonium were produced through the nitrate-reducing pathway [16]. Because of the well-known stable chemical property under the anoxic condition, ammonium will be accumulated in the formation water. Thus, the high concentration of ammonia was detected frequently in the production water from the oilfield, which is a problem in production water treatment for recycled water flooding and potentially imposed an environmental hazard. However, the detected concentration of ammonium was lower than the estimated value of stoichiometric calculation. To explain the elimination of ammonium in in situ oil reservoirs, we hypothesize that the anaerobic ammonium-oxidation (anammox) process might be responsible for consuming the ammonia in the anoxic oil strata. Because the anammox process was implicated to be most importantly responsible for the nitrogen cycle in all the anoxic habitats [11, 24, 31] and consistent with the growth condition of anammox bacteria, the subsurface oil reservoir formed a suitable niche for them.

The anammox bacteria are among the latest additions to the biogeochemical nitrogen cycle with ability to convert ammonium and nitrite into dinitrogen gas in a strictly anoxic environment. So far, five anammox genera have been identified with similar physiological property belonging to the Brocadiales order, while the phylogenetic distance is quite large with 16S rRNA gene sequence identities ranging from 87% to 99% [25]. Up to now, the presence of anammox bacteria has been demonstrated in worldwide oxygen-limited marine and freshwater systems and in the activated sludge [26, 27, 35, 47, 53, 54, 59], marine sediment and oxygen minimum zones [6, 9, 31, 32, 45, 55, 64], freshwater [22, 65], and river estuary ecosystem [8, 10]. Moreover, the phylogenetic biomarkers of the functional gene hzo and nirS encoding the key enzymes involving the anammox process were developed to investigated the functional diversity of anammox bacteria in wastewater treatment plants and natural environment [30, 32, 35, 47, 54, 57, 59], which significantly improved the understanding of the anammox bacterial community. However, the existence of anammox process in oil reservoirs is still unknown, and the knowledge of anammox in thermophilic habitats is also very limited. Given that the anammox bacteria were cultured in the experimental trail with optimum temperature below 43°C [58], the distribution of anammox bacteria in thermophilic habitats looks unfeasible. More recently, the discovery of anammox bacteria in the deep-sea hydrothermal vent and terrestrial hot springs substantially extends the temperature limit of anammox bacterial distribution [6, 22] by proving the presence and activity of anammox bacteria in the natural ecosystem with temperatures up to 100°C. The geothermal environment of an underground oil reservoir is a special hot habitat, while very little is known about the anammox bacterial distribution. Therefore, we initiated this survey of high-temperature oil reservoir ecosystems in order to find indications of other anammox bacteria that might thrive in the thermophilic environment and provide further information on bacterial nitrogen cycle in oil reservoirs. In this study, various oil reservoir samples were collected in Shengli Oilfield and observed by chemical and molecular ecology methods (i.e., 16S rRNA gene and hzo phylogeny). The existence of anammox bacteria was detected in the hot habitat to develop a general view of ecology and biogeochemistry characteristics of the anammox bacterial population in the geothermal and anoxic subsurface petroleum reservoir.

Materials and Methods

Sample Collection and Physicochemical Analyses

Shengli Oilfield is the second largest in China, located at Yellow River Delta in Shandong province in Eastern China. In total, 17 sampling wells distributed in an area of at least 2,000 km² were selected to represent contrasting geographical and biogeochemical attributes and oil production operation. Seven wells (S3–S9) were in the middle of the Yellow River Delta and two wells (S1, S2) in the vicinity of Bohai Sea (Fig. 1). The formation-water samples were collected directly from the well head in sterilized steel screw-cap bottles after flushing the lines for 30 min. The bottles were filled with oil/water mixture, sealed, and immediately stored in a large plastic box with ice and transported to the laboratory. The residual oil was removed by heating the sample to 50°C for 15 min and by phase separation in 2 l sterilized separatory funnel. Microbial biomass was collected from 2 l water phase by centrifugation at 15,000×g, 4°C. The chemical properties of the formation water were measured using Hach kits by following the manufacturer’s instructions (Hach, USA).

Generation of 16S rRNA Gene and hzo Clone Library

Total community DNA was extracted from cell pellets using the lysozyme/protease K/sodium dodecyl sulfate treatment followed by standard phenol/chloroform extractions. Nucleic acids were purified by the DNA purification kit (Promega, USA). Sample DNA was extracted several times, pooled, and concentrated. Polymerase chain reaction (PCR) amplification of 16S rRNA gene was performed by two protocols. Firstly, the initial PCR amplification was performed with forward primer Pla46F
in combination with the reverse primer 1037R [43], and the nested PCR was performed with the anammox-specific primers Amx 368F [3, 53] and Amx 820R [3, 52] according to the previous study [8]. Secondly, PCR was performed with the primer sets Brod541F [45] in combination with Amx820R [52] according to the previous study [23, 56]. The hzo gene fragments were amplified according to the previous study with two sets of primers hzocl1F1 and hzocl1R2 [54], and Ana-hzo1F and Ana-hzo2F [47]. PCR products from different primer sets were purified by gel cut procedure, and a combined library for the same sample with pMD-18T-vector (Takara, Japan) was constructed.

Sequencing and Phylogenetic Analysis

The insertion of 16S rRNA gene and hzo fragments was screened and retrieved by PCR amplification with the primer set of M13F and M13R. The different numbers of positive clones in each library were randomly selected for sequencing on an automated ABI Prism 3730 DNA analyzer by using M13 universal sequencing primers and the Big Dye Terminate kit (Applied Sciences, Foster City, CA). The sequences without chimeras were initially submitted to the FASTA program (version 3) [44] to determine their closest phylogenetic relatives. For 16S rRNA gene, DNA sequences were manually compiled and aligned using CLUSTAL X [62]. For hzo gene, nucleic acid sequences were first translated into amino acids, and the resulting protein sequences were aligned to their nearest neighbor using CLUSTAL X. Phylogenetic trees were constructed based on the Kimura two-parameter model [29] and the neighbor-joining algorithm [49] using the MEGA 4.0 software [60]. Bootstrap analysis with 1,000 replicates was applied to assign confidence levels to the nodes of the consensus trees.

Statistical Analyses

Operational taxonomic units (OTUs) for 16S rRNA and hzo gene diversity analysis were defined by 3% differences in nucleotide and amino acid sequences, as determined by using the furthest neighbor algorithm in DOTUR program [50]. DOTUR was also used to generate Chao, Shannon, and Simpson index for each clone library. The coverage of clone libraries was calculated from the equation $C = \frac{1}{\frac{1}{n} - \frac{1}{N} - \frac{1}{100}}$ [13]. Community classification of the anammox assemblages was determined with principal coordinates analyses (PCoA) [37]. Correlations between the anammox bacterial communities and environmental factors were determined by the canonical correspondence analysis (CCA) using the software CANOCO (ver. 4.5, Microcomputer Power, Ithaca, NY, USA) [61].

Real-Time PCR Assay

Gene copy numbers of total bacterial and anammox bacterial 16S rRNA gene in each sample were quantified with the quantitative PCR (qPCR) analysis. Real-time PCR was performed by using a FastStart Universal SYBR Green Master PCR mix (Roche Applied Science, Mannheim, Germany) in 96-well optical plates placed in an ABI 7000 Sequence detection system (Applied Biosystems, Foster City, CA). qPCR of total bacteria was performed with the primers 8F [2] and 338R [36] under the condition previously described [12, 35], and the qPCR assay of anammox bacteria was performed with the primers AMX-808-F and AMX-1040-R [17] under the adjusted condition
previously described [35]. Each reaction was performed in a 25-μl volume containing 1 μl of DNA template, 2.5 μl bovine serum albumin (0.1%), 0.5 μl of each primer (20 μM), and 12.5 μl of the ready reaction mix. The PCR cycle was started with 2 min at 50°C, 3 min at 95°C, followed by 45 cycles of 95°C for 30 s, annealing for 30 s at 55°C, and extension for 30 s at 72°C [35]. Cycle thresholds were determined by comparison with standard curves constructed using several concentrations of positive clone. The plasmid DNA concentration was determined on a Biophotometer (Eppendorf, Germany), and the copy number of target gene was calculated directly from the concentration of the extracted plasmid DNA. Tenfold serial dilutions of a known copy number of the plasmid DNA were prepared to generate an external standard curve. The \( R^2 \) values were greater than 0.98 for the curves. Samples and standards were prepared in triplicate. The specificity of the PCR amplification was determined by the melting curve and gel electrophoresis.

Nucleotide Sequence Accession Numbers

The GenBank accession numbers of 16S rRNA and \( hzo \) gene sequences are HM208755 to HM208788 and HM208789 to HM208820, respectively.

Results

Characteristics of Petroleum Reservoirs

The investigated oil reservoirs are between 55°C and 75°C and on average 1,200–1,900 m below the ground surface. The sampling wells experienced three different stages of crude oil recovery processes, namely primary production (S6, S7), secondary production (S1–S4, S8, and S9), and tertiary production (S5) (Table 1). Sampling wells S6 and S7 were of primary production containing water mainly from an aquifer in the strata. Wells S1–S5, S8, and S9 were developed for 5–10 years with water flooding to maintain the strata pressure and allow the extraction of residual oil. In particular, well S5 has been flooded for 2 years with injection of aqueous polyacrylamide solution. Overall, the formation water was carbonate type with slightly alkaline pH values between 7.3 and 8.1 under surface atmospheric pressure. The major physicochemical parameters for the geological formation water are presented in Table 1, showing low to moderate concentrations of sulfate, phosphate, and nitrate in the production water. The ammonium measured was high in the formation water (Table 1). Nitrite was near detection limit. Sulfide and ionic iron in S6 and S7 were higher than the other water samples.

<table>
<thead>
<tr>
<th>Number</th>
<th>Well number</th>
<th>Latitude (°N)</th>
<th>Longitude (°E)</th>
<th>Depth (m)</th>
<th>Temperature a (°C)</th>
<th>pH</th>
<th>WF (years)</th>
<th>CF (years)</th>
<th>Cont. (μg l⁻¹)</th>
<th>δS⁴ (ppm)</th>
<th>SO₄²⁻</th>
<th>PO₄³⁻</th>
<th>NO₃⁻</th>
<th>NH₄⁺</th>
<th>S²⁻</th>
<th>Fe²⁺</th>
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<td>7.7</td>
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<td>37.3</td>
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<td>0.06</td>
<td>0.06</td>
<td>0.02</td>
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WF: Water-flooding operation, CF: Chemical-flooding operation

*The average temperature of the oil-bearing horizons of the units.
Diversity of Anammox Bacteria in Petroleum Reservoirs

Seventeen samples with different production history from Shengli Oilfield were analyzed by PCR with anammox bacterial specific primer sets, but only nine samples were positive for anammox bacteria with the PCR primer sets used in this study. Subsequently, the nine samples were used to construct the 16S rRNA gene and hzo sequence libraries. Fifty to 80 clones of each gene library per sample were screened by PCR and then sequenced. The positive clones were obtained from 22 to 35 of 16S rRNA gene libraries and 23 to 34 of hzo gene libraries, with three to five and three to four OTUs, respectively (Table 2). The coverage value of all libraries ranged between 0.95 and 1.00, indicating that the majority of the phylotypes in the clone libraries was detected (Table 2). Based on both Shannon and Chaol estimator indices, the anammox bacterial community diversity of samples S3 and S4 was the highest, and that of samples S1 and S2 was the lowest, which was consistent with the property of S3 and S4 with the lowest temperature but S1 and S2 with the highest temperature.

Phylogenetic Affiliation of Anammox Bacteria in Oilfields

Anammox bacterial diversity was constructed with the community DNA obtained and then 16S rRNA gene clone library from the formation water or production water of the oil strata. Screening of 241 sequences without chimera against the FASTA database indicated that the clones matched sequences related to anammox bacterial genera: Candidatus Scalindua [53], Candidatus Kuenenia [59], Candidatus Brocadia [51], and Candidatus Jettenia [47]. Most of the sequence types had relatively high levels of similarity with their closest counterparts in public databases, whereas six phylotypes showed less than 92% sequence similarity to their nearest database entries and may belong to unknown phylotypes [48]. The phylogenetic relationships of the anammox bacteria-like 16S rDNA sequences from this study and their closest relatives are represented in a consensus tree (Fig. 2). Most of the OTUs fell into the Candidatus Scalindua cluster containing the species: Candidatus Scalindua brodae [53], Candidatus Scalindua sorokinii [31], and Candidatus Scalindua wagneri [53] with sequence identities of 90.1–99.6%, and this cluster was supported by 99% bootstrap value (Fig. 2). The sequences related to Candidatus sorokinii/brodae were similar to the sequence type Menez Gwen (AM941036), recovered from the hydrothermal deep-sea vent recently [6]. In addition, six phylotypes in this subcluster were further divided into two groups that are distantly affiliated with each other (Fig. 2). The remaining sequences were more diverse and distributed within the Candidatus Scalindua wagneri subcluster, such as the Colne Estuary (EU394279) in the UK [10] and the Cape Fear River Estuary (FJ490132) in North Carolina [8], which were isolated from freshwater systems. The second large fraction of sequences were grouped within Candidatus Kuenenia genus (Fig. 2); the most abundant phylotypes, S4E6 and S2B10, in this cluster were affiliated with the sequence obtained from California Leonard hot spring clone HRH850 from a 52°C spring [22] and Rainbow hydrothermal chimney clone (AM941023) which was the first finding of anammox bacteria in the hot habitat of an active deep-sea vent with environmental fluid temperatures up to 100°C [6]. In addition, the Brodcadia cluster included the clones from high-temperature wells S1,

### Table 2 Diversity and predicted richness of the anammox 16S rRNA gene and hzo sequences recovered from the production water samples of Shengli Oilfield

<table>
<thead>
<tr>
<th>Number</th>
<th>Well number</th>
<th>Number of sequences</th>
<th>Number of OTUs&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Coverage&lt;sup&gt;b&lt;/sup&gt;</th>
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<th>Chao1</th>
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<td>22</td>
<td>26</td>
<td>3</td>
<td>4</td>
<td>1.00</td>
</tr>
</tbody>
</table>

<sup>a</sup>Unique OTUs were determined using the DOTUR program based on the 3% sequences difference

<sup>b</sup>The coverage, Shannon-Weiner, Simpson and S<sub>Chao1</sub> richness estimators were calculated using the OTUs data
S2, and S8, which were closely related to the sequences of California Leonard hot spring clone HRH852 [22] and Geothermal stream clone HAuD-MB/2-35 from hot water above 50°C in a Japanese gold mine [19]. The other sequences belonging to Jettenia cluster were isolated from lower temperature oilfields, closely related to the sequences from freshwater environments (Fig. 2) of the Cape Fear River Estuary in the UK [8]. Interestingly, cluster A containing clones S6G9, S7G9, S1B7, and S2E8 were distantly (90.1–92.6%) related to the known anammox
bacteria, while this cluster was adjacent to the Scalindua cluster in the topology structure of the phylogenetic tree (Fig. 2), which were retrieved from the no-water-flooding and highest temperature oilfields, respectively. Samples S3, S4, S6, and S7 yielded four sequence types forming cluster B (Fig. 2), affiliated with the Rainbow hydrothermal chimney clone CH10MOM07.5 (AM941027) from the high-temperature deep-sea vent and the Nevada Patua-Hazen hot spring clone HRH860 from a 51°C spring [22], inferring a potentially new anammox bacterial cluster [6].

Functional Diversity of Anammox Bacteria by \textit{hzo} Gene Analysis

The HZO enzyme (hydrazine oxidoreductase) catalyzes the initial hydrazine-forming reaction during the anammox biochemical process. The encoding gene \textit{hzo} cluster 1 [54] was proven to be an appropriate and feasible biomarker for phylogenetic analysis of anammox bacteria in the enrichment cultures and environmental samples [34, 54]. To detect the functional diversity of anammox bacteria in the petroleum reservoirs, clone libraries of \textit{hzo} gene were constructed by using the community DNA from the formation-water samples based on the cluster 1 \textit{hzo} specific primers sets. The deduced amino acid sequences of 243 nucleic acid sequences could be grouped into 12 OTUs based on 3% substitutions/difference of amino acid sequences. The comparative analysis indicated that the closest matched HZO sequences belong to the anammox bacteria \textit{Candidatus Anammoxoglobus}, \textit{Candidatus Kuenenia}, \textit{Candidatus Jettenia}, and \textit{Candidatus Scalindua} (Fig. 3), but no sequence type fell into the anammox bacteria \textit{Candidatus
Brocadia. The phylogenetic relationships of the HZO-like sequences in this study and their closest relatives were represented in a consensus tree, and the amino acid sequences of HZO cluster 2a [54] were used as out-groups to root the tree (Fig. 3). The most abundant clones were grouped in the Scalindua cluster, which was 96.5–98.9% similar to HZO sequences of Candidatus Scalindua sp. [54]. Adjacent to this cluster, cluster D was distantly related to other HZO phylotypes (Fig. 3). Consistent with cluster C of 16S rRNA gene tree, this cluster had the same phylogenetic topology status in the HZO tree, so it was likely that sequences of these two clusters represented the unknown anammox bacteria. Cluster D sequences also formed a slightly distant group affiliated with other detected hzo genes. However, no corresponding cluster sequences to cluster C were found in the 16S rRNA gene libraries (Figs. 2 and 3). Most sequences from water-flooding and lower temperature oilfields were grouped within Jettienia cluster, showing 97.6–99.5% sequence identities to the HZO sequence of Candidatus Jettienia asiatica [47]. But some of no-water-flooding oilfield sequences fell into Anammoxoglobus cluster with 98.2–99.6% identities to the HZO sequence of Candidatus Anammoxoglobus sp. [54]. Consistent with the 16S rRNA gene phylogeny, most of the highest temperature oilfield sequence types were related to the HZO sequence of Candidatus Kuenenia stuttgartiensis with 97.3–99.8% sequence identities [59]. Interestingly, no sequences related to Candidatus Brocadia were detected in any of the samples used in this study.

Quantification of Anammox Bacteria

The abundance of total bacteria and anammox bacteria in the oilfield production water was estimated based on the quantification of bacterial 16S rRNA genes using the qPCR method [12, 17, 35]. The amount of total bacterial and anammox bacterial 16S rRNA gene copies ranged from 2.3±0.18×10^4 to 7.3±0.65×10^6 ml^-1 and from 6.6±0.51×10^2 to 4.9±0.36×10^3 (Fig. 4), respectively. Based on the estimation that 3.6 copy numbers of 16S rRNA gene per bacterial cell genome [18], the total bacterial density was calculated from 6.4±0.5×10^3 to 2.0±0.18×10^6 cells ml^-1. Assuming that the anammox bacteria contained one copy of the 16S rRNA operon based on Candidatus Kuenenia stuttgartiensis genome [59], the anammox bacterial density was calculated to be 6.6±0.51×10^2 to 4.9±0.36×10^3 cells ml^-1. The anammox bacteria was a small fraction of the bacterial community in Shengli Oilfield production waters, as the anammox bacteria accounted for 2.5% to 10.3% of all bacteria detected. The highest concentration of total bacteria and anammox bacteria was found in the production water from S4 (Fig. 4), where the lowest temperature and moderate time scale of water-flooding operation were observed (Table 1). Correlation analysis showed that the total bacteria and anammox bacteria populations were highly related to the temperature gradient. The lowest bacterial abundance was found in sample S2, where the highest temperature and longest term of flooding were observed.

Anammox Bacterial Communities Classification

Significant difference of the anammox bacterial assemblages in the oilfields was revealed by the PCoA test (p<0.005) [38] based on the phylogenetic tree of 16S rRNA gene and HZO protein sequences. The results indicated that anammox bacterial assemblage fell into four groups, S5 for chemical flooding, S3 and S4 for lower temperature and water-flooding oilfield, S6 and S7 for no-water-flooding oilfield, and S1, S2, S8, and S9 for high-temperature water-flooding oilfield (p<0.032) (Fig. 5a). Both S6 and S7 represented no-water-flooding oil reservoirs, and S5 represented chemical-flooding oilfield, showing obvious difference from the other samples from the water-flooding oilfield along the first principal coordinate (P1) which explains 32.9% of the total anammox bacterial community variability among all the sampling oilfields (Fig. 5a). The PCoA analysis of the 16S rRNA gene and HZO protein sequences of anammox bacterial assemblages showed a slightly different classification (Fig. 5b), with S9 closer to S8 in the former analysis, but S8 was closer to S2 in the latter one.

Spatial Distribution of Anammox Bacterial Communities

To find the correlations between the distribution of anammox bacterial community and the environmental variables of the oilfields, canonical correspondence analysis was conducted based on the 16S rRNA gene and HZO
protein sequences and the physiochemical parameters recovered in Shengli Oilfield. The results showed that the CCA axe in 16S rRNA gene deduced scheme explained 40.1% but the CCA axe in HZO sequence deduced scheme explained 54.6% of the cumulative variance of the correlation between the environmental factors and the anammox bacterial community distribution (Fig. 6). The CCA analysis indicated that all of the anammox bacterial assemblages fell into four groups, with the CCA1 distinguishing the anammox bacterial assemblages of S1, S2, S8 and S9, S6, and S7 together and S5 distant to others, while the CCA2 axe found S3 and S4 as one assemblage (Fig. 6). The oil recovery process contributed another important effect ($p=0.005$, 1,000 Monte Carlo permutations) to the distinction of the anammox bacterial community and environment relationship, which revealed that the distribution of the anammox bacterial assemblages of S1, S2, S8, and S9 was positively correlated with the water flooding but in the opposite direction to the distribution of S6 and S7, and the distribution of S5 was positively correlated with chemical flooding (Fig. 6). The temperature and the concentration of ammonium cation appeared to be significant ($p=0.048$, 1,000 Monte Carlo permutations) to the anammox bacterial distribution, and the environmental parameter relationship

**Figure 5** PCoA ordination diagrams of the anammox bacterial assemblages calculated with 16S rRNA gene sequences (a) and HZO protein sequences (b) from production water of Shengli Oilfield. Shown are the plots of the first two principal coordinate axes (P1 and P2) for PCoA and the distributions of the anammox bacterial assemblages (designated with the sampling wells) in response to these axes.

**Figure 6** CCA ordination plots for the first two dimensions to show the relationship between the anammox bacterial diversity with environmental parameters analyzed using a 16S rRNA gene sequence and b HZO protein sequence in the production water of Shengli Oilfield. Correlations between environmental variables and CCA axes are represented by the length and angle of arrows (environmental factor vectors). *Temp* temperature, *WF* water flooding, *CF* chemical flooding.
with the S1, S2, S8, and S9 were positively correlated with the two factors. Other environmental factors did not contribute significantly (p > 0.500) to the anammox bacterial community, while the combination of these environmental factors provided additionally 55.0% of the total CCA explanatory power.

Discussion

The combined data of clone libraries of both 16S rRNA and hzo genes confirmed the occurrence of anammox bacteria in the anoxic geothermal oilfield environment at Shengli Oilfield in China. Our results confirmed that Candidatus Brocadia sp. and Candidatus Kuenenia sp. could be detected in oil reservoirs with temperature up to 75°C and were the first to detect the anammox bacterial species Candidatus “Scalindua sinooilfield” in such habitat. It is consistent with other findings observed in the deep-sea hydrothermal vent at Mid-Atlantic Ridge [6] and in Great Basin hot springs of Northern California and Western Nevada [22]. By applying more specific primer sets for anammox bacterial 16S rRNA gene and the functional biomarker hzo gene, anammox bacteria were successfully detected in the oil reservoir ecosystem. This is the first report for the existence of anammox bacteria in the high-temperature oil reservoir. Therefore, anammox bacteria are present not only in the marine environments at generally lower ambient temperatures [55, 64] but also in the high-temperature environment, which expands our knowledge of the biogeography of anammox bacteria.

The diversity of anammox bacteria depends on the oil production processes and physicochemical conditions in situ. Most phylotypes detected in relatively higher temperature wells belong to Kuenenia and Brocadia clusters. Although the highest optimum temperature was at 43°C in enrichment culture [58], Candidatus Kuenenia species were also obtained from the hydrothermal vent with fluid temperatures more than 100°C and the hot springs with higher than 52°C [6, 22]. In this study, the Brocadia-like organisms were detected in the water samples from wells S1, S2, and S8 with temperature up to 75°C, which was the first report of the Brocadia-like organisms in such habitat. Interestingly, the sequence detected in the no-water-flooding wells belonged to the Anammoxoglobus cluster closely related to Candidatus “Anammoxoglobus propionicus” which had capability to oxidize simple organic acids (acetate and propionate) at a higher rate and without a transient nitrite accumulation [27]. This property may be critical for them to thrive in an oil reservoir environment because the crude oil consists of abundant hydrocarbons which can be converted to short-chain fatty acids by microbial fermentation process in the oil reservoirs under primary operation without anthropogenic influences, such as water flooding or chemical injection [14, 15].

For a more comprehensive analysis of the functional diversity of anammox bacteria in oil reservoirs, the hzo sequences were retrieved to compare the phylogenetic diversity of anammox bacterial 16S rRNA gene. Consistent with previous results [34, 35, 54], the topologies of the phylogenetic tree of the obtained hzo was congruent with retrieved 16S rRNA gene phylogeny, suggesting that the hzo gene is a good alternative biomarker for detecting the anammox bacteria in oil reservoirs. More importantly, our results indicated that the phylogenetic diversities based on 16S rRNA and hzo genes are complementary to each other for Brocadia-like organisms in the 16S rRNA gene library and Anammoxoglobus-like ones in hzo library. Besides the sequences closely related to the known anammox bacteria, two clusters of sequences retrieved were apparently more distantly related to the sequences from cultured and uncultured anammox bacteria. Firstly, cluster A 16S rRNA gene sequences were distantly related to other Candidatus Scalindua-like phylotypes. Based on the sequence identities of up to 97% of the 16S rRNA gene commonly used to define different species [48], it was proposed that the sequences of this cluster comprise a new species within the genus Candidatus Scalindua, for which the name Candidatus “Scalindua sinooilfield” is proposed provisionally. Given the cluster D hzo sequences have similar topology structure compared to the cluster A 16S rRNA gene phylogeny, we suggest that this cluster belongs to the new anammox bacterial genus Candidatus “Scalindua sinooilfield”. Assuming this hypothesis is correct, the distinctness of the Candidatus “Scalindua sinooilfield” cluster could also explain that cluster D hzo sequences have distinctly different lengths compared to other hzo genes of Candidatus Scalindua species. In addition, the cluster C hzo sequences show further separation from other hzo-like sequences, whereas the corresponding sequence cluster could not be found in the 16S rRNA gene phylogenetic tree, which might be due to the differences of specificity and efficiency of the PCR primer sets of these two biomarkers [34]. Thus, to further investigate the functional diversity of anammox bacteria in various ecosystems, more effective primers need to be designed for the hzo or other important anammox genes, such as the putative cytochrome cd1-containing nitrite reductase gene (nirS) proven to be an alternative biomarker for Scalindua-like sequences recently [32].

The abundance of anammox bacteria decreased in the oil reservoirs as the temperature increased. The lowest concentration was detected in samples with moderate water flooding, which was comparable with the anammox bacterial numbers in high-temperature deep-sea vents [6]. However, the significant variation of anammox bacterial
abundance was observed in different oilfields with temperatures ranging from 55°C to 75°C. The mean concentration of anammox bacteria was obviously lower than that in marine ecosystem [8, 17, 55]. This discrepancy might be related to the inhibition of high temperature to the growth of anammox bacteria. Meanwhile, the water-flooding and chemical-flooding operations seem to affect their number. The anammox bacterial density in the moderate water-flooding oilfield was higher than that in the no-water-flooding and the longest water flooding. The chemical-flooding oilfield S5 was lower by 90 times than that in the no-chemical-flooding oilfields. The abundance of anammox bacteria in primary production well samples was between those of the chemical-flooding sample and the water-flooding samples.

The distribution of anammox bacteria is affected by the oilfield history and production processes. Both temperature and recovery process are the two most important environmental variables affecting the distribution of anammox bacteria in the oil reservoirs. Most phylotypes detected in primary production wells belong to Scalindua cluster observed from the freshwater system, which might adapt to the oil reservoir environment in underground water system, while most phylotypes in the water-flooding and chemical-flooding oil reservoir samples fall into the Jettenia cluster, which might be originated from the injected water. This discrepancy might be related to the introduction of microorganisms by injected water or chemical [7, 33], resulting in the shift of microbial community composition. Given that the members of Candidatus Scalindua and Candidatus Jettenia genera were previously proved to be ubiquitous distributed in environments such as river and marine environments or granular sludge anammox reactor [47, 55, 64, 65], it is reasonable to have anammox bacteria in the injection water with sources of river water or recycled flooding water. This explains the existence of the putative Scalindua and Jettenia-related sequences in the production water from oilfield. Interestingly, there are more than one kind of anammox bacteria detected in most of the oil reservoirs, which contradict the niche differentiation of anammox bacteria, genera Candidatus Kueneinia, Candidatus Anamoxoglobus, and Candidatus Brocadia in anoxic freshwater and Candidatus Scalindua in anoxic marine habitats [55], indicating that anammox bacteria have broader distribution than previously estimated [27, 55].

The existence of anaerobic ammonium-oxidizing bacteria in the high-temperature oil reservoir was confirmed in this study. Widely distributed and diverse 16S rRNA gene and hzo sequence types were detected in the production water samples. In addition, two clusters of sequences retrieved from high-temperature oilfield might represent a novel anammox bacterial phylotype. Anammox bacterial communities are influenced by temperature and production processes. Further study of anammox bacteria in oilfield ecosystems would allow a better understanding of the nitrogen cycle in the subterranean oil reservoir ecosystems.

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