Authors
Jenna L Shelton, Denise M Akob, Jennifer C McIntosh, Noah Fierer, John R Spear, Peter D Warwick, and John E McCray

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Environmental Drivers of Differences in Microbial Community Structure in Crude Oil Reservoirs across a Methanogenic Gradient

Jenna L. Shelton 1*, Denise M. Akob 2, Jennifer C. McIntosh 1,3, Noah Fierer 4,5, John R. Spear 6, Peter D. Warwick 1 and John E. McCray 6,7

1 Eastern Energy Resources Science Center, U.S. Geological Survey, Reston, VA, USA, 2 National Research Program-Eastern Branch, U.S. Geological Survey, Reston, VA, USA, 3 Department of Hydrology and Atmospheric Sciences, University of Arizona, Tucson, AZ, USA, 4 Department of Ecology and Evolutionary Biology, University of Colorado, Boulder, CO, USA, 5 Cooperative Institute for Research in Environmental Science, University of Colorado, Boulder, CO, USA, 6 Department of Civil and Environmental Engineering, Colorado School of Mines, Golden, CO, USA, 7 Hydrologic Science and Engineering Program, Colorado School of Mines, Golden, CO, USA

Stimulating in situ microbial communities in oil reservoirs to produce natural gas is a potentially viable strategy for recovering additional fossil fuel resources following traditional recovery operations. Little is known about what geochemical parameters drive microbial population dynamics in biodegraded, methanogenic oil reservoirs. We investigated if microbial community structure was significantly impacted by the extent of crude oil biodegradation, extent of biogenic methane production, and formation water chemistry. Twenty-two oil production wells from north central Louisiana, USA, were sampled for analysis of microbial community structure and fluid geochemistry. Archaea were the dominant microbial community in the majority of the wells sampled. Methanogens, including hydrogenotrophic and methylotrophic organisms, were numerically dominant in every well, accounting for, on average, over 98% of the total Archaea present. The dominant Bacteria groups were Pseudomonas, Acinetobacter, Enterobacteriaceae, and Clostridiales, which have also been identified in other microbially-altered oil reservoirs. Comparing microbial community structure to fluid (gas, water, and oil) geochemistry revealed that the relative extent of biodegradation, salinity, and spatial location were the major drivers of microbial diversity. Methanogens, including hydrogenotrophic and methylotrophic organisms, were numerically dominant in every well, accounting for, on average, over 98% of the total Archaea present. The dominant Bacteria groups were Pseudomonas, Acinetobacter, Enterobacteriaceae, and Clostridiales, which have also been identified in other microbially-altered oil reservoirs. Comparing microbial community structure to fluid (gas, water, and oil) geochemistry revealed that the relative extent of biodegradation, salinity, and spatial location were the major drivers of microbial diversity. Archaela relative abundance was independent of the extent of methanogenesis, but closely correlated to the extent of crude oil biodegradation; therefore, microbial community structure is likely not a good sole predictor of methanogenic activity, but may predict the extent of crude oil biodegradation. However, when the shallow, highly biodegraded, low salinity wells were excluded from the statistical analysis, no environmental parameters could explain the differences in microbial community structure. This suggests that the microbial community structure of the 5 shallow, up-dip wells was different than the 17 deeper, down-dip wells. Also, the 17 down-dip wells had statistically similar microbial communities despite
**INTRODUCTION**

Increasing demands for energy combined with diminishing economically accessible fossil fuel reserves will require novel energy-efficient technologies for hydrocarbon production. There are large, historically non-viable estimated global reserves of both heavily biodegraded (i.e., microbially altered) crude oil and residual crude oil (crude oil that remains in situ following traditional extraction mechanisms). However, these reservoirs are increasingly being considered a potentially viable contribution to the energy mix as lighter, easier-to-produce crude oils diminish (Hein et al., 2013). One suggested mechanism to "produce" this difficult-to-obtain oil is through enhancing in situ microbial communities to metabolize the oil to natural gas via methanogenic crude oil biodegradation (Parkes, 1999; Gieg et al., 2008; Jones et al., 2008; Head et al., 2014; Meslè et al., 2015). However, significant technical barriers to commercial development remain, including identification and manipulation of the most important biogeochemical factors controlling bioconversion.

A consortium of microorganisms, most notably, methanogens and syntrophic bacteria, perform methanogenic crude oil biodegradation by breaking down crude oil into methanogenic substrates (e.g., formate, acetate, carbon dioxide), which are subsequently converted into natural gas. The process has been confirmed in laboratory studies where microbial communities collected from deep methanogenic oil reservoirs can consume crude oil and produce methane (e.g., Gieg et al., 2008; Berdugo-Clavijo et al., 2012; Berdugo-Clavijo and Gieg, 2014). In addition, subsurface communities capable of converting oil to natural gas have been described using a variety of sequencing techniques, but community structure (abundance and diversity) varied between studies (e.g., Dahle et al., 2008; Pham et al., 2009; Shartau et al., 2010; Yamane et al., 2011; Kryachko et al., 2012; Li et al., 2012; Berdugo-Clavijo and Gieg, 2014; Meslè et al., 2015). Despite this knowledge, we are still limited in our understanding of how hydrochemistry and microbial populations may impact methanogenic crude oil biodegradation under fairly constant lithological conditions (e.g., Kirk et al., 2015). Most of the previous studies were performed on a small scale (i.e., a small number of samples from a hydrogeochemically similar setting), while the few large datasets rarely compare microbial communities across environmental gradients. In addition, it is difficult to analyze the impact of hydrogeochemical conditions across samples collected from varying lithologies, because lithology has a causative impact on hydrochemistry.

The objective of this study was to expand on current knowledge about the microbiology of methanogenic crude oil biodegradation by sampling biomass across a hydrogeochemical gradient within the same lithologic formation exhibiting varying extents of methanogenesis and crude oil biodegradation. To accomplish this objective, we characterized microbial communities from 22 different oil wells from 6 different oil fields across a transect; all wells produce from the Wilcox Group of north-central Louisiana, USA. Previous work (Warwick et al., 2008; McIntosh et al., 2010; Shelton et al., 2014, 2016) established the amount of crude oil biodegradation, relative amount of microbial methanogenesis, and the formation gas and water geochemistry across north-central Louisiana. The new microbial data, combined with previous analyses of fluid chemistry, enabled us to explore the environmental conditions and microbial populations most conducive to methanogenic crude oil biodegradation.

**STUDY SITE AND METHODS**

**Study Site**

Twenty-two oil wells (Table S1), spanning 6 oil fields across an approximately 65 km transect producing from the Paleocene–Eocene Wilcox Group were sampled in 2014 (Figure 1). Biomass, produced water, and 4 oil samples were collected in this 2014 sampling campaign. These data were supplemented by produced gas and oil data collected from a 2011/2012 sampling campaign (Shelton et al., 2014, 2016). We justified the use of this complimentary, older dataset (oil and gas data from 2011/2012) due to slow rates of crude oil biodegradation and subsequent methanogenesis (e.g., Larter et al., 2003, 2006), and because chemical water analyses from this sampling campaign produced values similar to those from 2011/2012 (Shelton et al., 2014). The sampled wells spanned a wide range of geochemical parameters, including temperature, pH, and salinity (Table 1), and formation fluids with varying extents of both crude oil biodegradation and microbial methanogenesis (McIntosh et al., 2010; Shelton et al., 2014, 2016). One oil field sampled, the Olla Field, has been previously suggested to be a hot spot of microbial methanogenesis in the region (McIntosh et al., 2010), later attributed to ideal geochemical conditions for microbial methanogenesis (Shelton et al., 2014).

**Field and Analytical Methods**

Formation oil, water, and gas were collected over two different sampling campaigns in August 2011, and July 2012, while biomass and formation water and oil were collected in August 2014 (Table S1). Due to the similarity of the produced water geochemistry collected in 2011/2012 compared to 2014, along with slow rate constants for crude oil biodegradation (e.g., Larter
et al., 2006), major changes in methanogenic activity and crude oil composition were not expected over the sampled time scale (2–3 years). Each well is identified by oil field and a sample number (Table 1, Table S1). The up-dip, shallow (393–472 m) wells from the Crossroads, Salt, and Tullos Urania fields are identified as CR, S, or TU wells, respectively. The mid-depth (667–1044 m), centrally-located wells from the Olla and Nebo-Hemphill fields are identified as O or NH, respectively. The deep (1063–1588 m), down-dip wells from the Cypress-Bayou Field are identified as CB.

Formation water and oil were collected directly from the wellhead into a 5 gallon Nalgene carboy, where the temperature of the brine/oil mixture was taken while the mixture partitioned, using an Oakton Acorn Temp 6 Thermometer (Vernon Hills, Illinois USA). Once separated, the water was released from a spigot at the bottom of the carboy into sterile 60 mL BD Luer-Lok syringes, and filtered through attached 0.2 µm nylon syringe filters into 30 mL HDPE bottles. The HDPE bottles were pre-cleaned and the collected fluids were preserved according to associated analysis (e.g., Shelton et al., 2014), filled with no headspace, capped, put on ice, and shipped to either the University of Arizona or the U.S. Geological Survey (USGS) where they were kept at 4°C until analyzed.

Produced fluids were collected in 2 L pre-combusted, sterile glass bottles for filtering microbial biomass. Nalgene tubing was inserted into the bottle after the oil/brine mixture was allowed to partition, and the partitioned brine was filtered through sterile 0.22 µm Sterivex™ GP filter units (Millipore®, Billerica, MA USA) using a GeoPump™ (Geotech Environmental Equipment, Inc. Denver, CO). The volume of filtered water was recorded, and triplicate filters were taken for each well. Filters were immediately frozen on dry ice and shipped to Colorado School of Mines, where they were kept at −80°C until DNA extraction. Alkalinity was titrated within 8 h of sample collection following the protocol outlined in Gieskes and Rogers (1973), and pH was recorded in the field using a Thermo Scientific Orion pH electrode. The δ13C-DIC of produced water was measured at the University of Arizona’s Environmental Isotope Laboratory using a ThermoQuest Finnigan Delta Plus XL continuous flow gas ratio mass spectrometer (precision at least ±0.3‰). Anions
TABLE 1 | Major field and geochemical parameters from this study and others and associated bins for statistical analyses.

| Well ID | Oil Field | Depth (m) | Temp. °C | pH | δ¹³C-DIC ‰ | Alk. | Cl-mM | δ¹³C-CO₂ ‰ | δ¹³C-CH₄ ‰ | Extent of Meth. | Extent of COB | CH₄ Prod. *m³ CH₄/day |
|---------|-----------|-----------|----------|----|-------------|------|-------|-------------|-------------|----------------|--------------|---------------------|
| CR1     | Crossroads| 438.3     | 28.7     | 7.6| −2.2        | 13.4 | 567   | n/a         | n/a          | Low           | High         | BDL                 |
| CR2     | Crossroads| 439.8     | 25.5     | 7.7| −2.0        | 13.2 | 559   | n/a         | n/a          | Low           | High         | BDL                 |
| CR3     | Crossroads| 472.4     | 24.2     | 7.6| −2.0        | 13.2 | 564   | n/a         | n/a          | Low           | High         | BDL                 |
| S3      | Salt      | 433.7     | 28.4     | 7.5| −0.3        | 10.2 | 685   | n/a         | n/a          | Low           | High         | BDL                 |
| TU1     | Tullos Urania| 383.2     | 32.0     | 7.2| 2.9         | 7.4  | 776   | n/a         | n/a          | Low           | High         | BDL                 |
| O3      | Olla      | 828.4     | 39.8     | 7.1| 16.9        | 37.7 | 1382  | 12.8        | −52.3        | High          | Minimal     | 590,000             |
| O4      | Olla      | 850.4     | 41.7     | 6.7| 15.7        | 40.8 | 1394  | 10.9        | −45.8        | Medium        | Minimal     | 160,000             |
| O5      | Olla      | 833.3     | 38.0     | 7.0| 22.8        | 34.8 | 1334  | 17.2        | −45.3        | Medium        | Moderate    | 110,000             |
| O6      | Olla      | 853.7     | 43.2     | 6.7| 21.8        | 45.6 | 1394  | 16.7        | −46.3        | High          | Moderate    | 390,000             |
| O7      | Olla      | 850.4     | 41.8     | 7.4| 21.5        | 48.2 | 1371  | 16.2        | −46.9        | High          | Moderate    | 390,000             |
| O8      | Olla      | 856.5     | 40.3     | 7.1| 23.6        | 52.2 | 1351  | 17.7        | −49.7        | High          | Minimal     | 460,000             |
| O18     | Olla      | 833.9     | 38.4     | 6.9| 21.2        | 44.1 | 1385  | 15.8        | −47.6        | Medium        | Moderate    | 130,000             |
| O26     | Olla      | 679.4     | 33.7     | 6.9| 23.4        | 38.2 | 1154  | 15.8        | −48.3        | High          | Minimal     | 430,000             |
| O27     | Olla      | 666.9     | 33.6     | 7.1| 23.6        | 51.8 | 1117  | 17.4        | −45.3        | High          | Minimal     | 190,000             |
| O31     | Olla      | 1043.8    | 48.5     | 6.9| 21.7        | 56.4 | 1678  | n/a         | n/a          | High          | n/a         | n/a                 |
| O32     | Olla      | 856.5     | 40.3     | 6.9| 24.1        | 53.1 | 1385  | n/a         | n/a          | High          | n/a         | n/a                 |
| NH1     | Nebo-Hemphill| 1208.2    | 44.7     | 6.9| 8.6         | 5.8  | 1676  | 1.8         | −69.0        | Low           | Moderate    | n/a                 |
| NH2     | Nebo-Hemphill| 1062.5    | 45.0     | 6.9| 16.0        | 10.3 | 1512  | n/a         | −48.9        | Medium        | n/a         | n/a                 |
| NH3     | Nebo-Hemphill| 1211.9    | 38.0     | 6.8| 8.3         | 6.7  | 1687  | 2.2         | −56.4        | Low           | Minimal    | 25,000               |
| CB1     | Cypress-Bayou| 1472.6    | 24.4     | 6.8| 1.2         | 5.0  | 2020  | −6.3        | −62.7        | Low           | Minimal    | BDL                 |
| CB3     | Cypress-Bayou| 1485.5    | 29.4     | 6.9| 0.2         | 4.7  | 2066  | −6.8        | −63.2        | Low           | Minimal    | BDL                 |
| CB4     | Cypress-Bayou| 1588.0    | 28.8     | 7.3| 0.7         | 5.6  | 2000  | n/a         | n/a          | Low           | Minimal    | BDL                 |

*Shelton et al. (2014); Shelton et al. (2016). Fill color coordinates to the bin each sample value was placed in for statistical analysis. Red, Highest; Orange, High; Tan, Medium; Light Blue, Low; Blue, Lowest. Cells labeled “n/a” (white) were not binned.

Temp., Temperature; DIC, Dissolved Inorganic Carbon; Alk., Alkalinity; CO₂, Carbon Dioxide; CH₄, Methane Gas; Meth., Methanogenesis; COB, Crude Oil Biodegradation; BDL, Below Detection Limit; CH₄ Prod., methane produced from the wellhead.

n/a Indicates that the sample(s) were not available for analysis.

Microbial Community Analysis

DNA extractions, amplifications, and Illumina MiSeq 16S rRNA gene sequencing were performed at the University of Colorado Next Generation Sequencing Facility. DNA was extracted using a MO BIO PowerSoil® DNA Isolation Kit (MO BIO Laboratories, Carlsbad, CA) following the manufacturer’s protocol, with slight modifications for the Sterivex™ filter units. The Sterivex filter units were opened and slices of the filter were added directly to the bead tubes. Extracted DNA was amplified using the 515-F (5’-GTGCCAGCMGCCGCGGTAA-3’) and 806-R (5’-GGACTACHVGGGTWTCTAAT-3’) 16S rRNA gene primer pair (Fierer et al., 2012); the primers also included Illumina adapters and reverse primers (adapted with error-correcting 12-bp barcodes unique to each sample). PCR was performed with a GoTaq® Hot Start PCR Master Mix (Promega, Madison, WI USA) in a 25 µL reaction, with thermal cycling consisting of initial denaturation at 94°C for 3 min, followed by 35 cycles (45 s) of denaturation at 94°C, annealing at 50°C for 30 s, extension at 70°C for 90 s, and a final extension at 72°C for 10 min. Successful amplification was verified with gel electrophoresis (2% agarose gel), and the amplified DNA was sequenced using the Illumina MiSeq platform running 2 x 250 bp chemistry (Illumina, San Diego, CA USA).

Downstream processing was performed using the UPARSE pipeline (Edgar, 2013), where forward reads were demultiplexed using an in-house custom Python script (https://github.com/leff/helper-code-for-uparse). Quality filtering was conducted using a maxee value of 0.5 (filtering out ca. 10% of the nucleotides). Prior to determining and assigning phylotypes, the sequences were dereplicated and singleton sequences were removed. Taxonomic units were mapped to the raw sequences at the 97% similarity threshold using the Greengenes 13.8 (http://greengenes.secondgenome.com) database. Any sequences that matched phylotypes representing chloroplasts or mitochondria were removed, along with sequences matching those found in the blank sample (i.e., sequencing contaminants). The samples were then rarefied randomly at 3700 sequences per sample using the biom package in R (McDonald et al., 2012; http://www.r-project.org). Sequence reads for each well were deposited to the National Center for Biotechnology Information Short Read
Microbial Community Statistics

The 22 sampled wells were binned according to each of the following environmental parameters in order to determine how changes in environmental parameters impacted microbial community structure: oil field and well depth; temperature, pH, chloride concentration, indicators of extent of methanogenesis (δ¹³C-DIC values, alkalinity, methane production volumes, δ¹³C-CO₂ values, and δ¹³C-CH₄ values), and extent of crude oil biodegradation (Table 1). Each sampled well was assigned to one of 3–5 categories—Lowest, Low, Medium, High, or Highest—for each environmental parameter analyzed for that sample. The thresholds for these bins were determined so that a relatively even number of samples (e.g., 5–6 samples in each of the 3–5 categories) were placed in each bin (Table 1). The binning process was aided by natural cut-offs for many of the environmental parameters (e.g., chloride concentrations increased from 776 to 1154 mM between two different oil fields). Extent of methanogenesis and extent of crude oil biodegradation rankings for each well were assigned using indicators of methanogenesis (e.g., δ¹³C-DIC values) and crude oil geochemistry (e.g., pristane/phytane values), respectively (see Shelton et al., 2016 for methodology). Any well with missing data for a specific geochemical analysis was omitted from any statistical analyses performed using that parameter. All statistical analyses were conducted using the vegan package (Oksanen et al., 2015) in R (http://www.r-project.org).

RESULTS AND DISCUSSION

Produced Fluid Geochemistry

Formation water chemistry was sampled in both 2011/2012 and for this study, but only the data for the current sampling campaign will be presented here. Formation water temperatures ranged from 22.4° to 48.5°C across the sampled transect, with the coldest water located in the Crossroads Field and the warmest water located in the Olla Field (Table 1). The pH ranged from 6.7 to 7.7, with the lowest pH found in the Olla Field and the highest pH found in the Crossroads Field. Alkalinity values ranged from 22.4 to 24.1‰, with the highest values found in the Olla and Nebo-Hemphill fields. The δ¹³C-CO₂ and δ¹³C-CH₄ values were taken from Shelton et al. (2014), as gas samples were not collected for this study. Daily gas production volumes were supplied by XTO Energy (written communication), and were converted to daily methane production volumes (Table 1) using gas composition data presented in Shelton et al. (2014).

Differences in Microbial Community Composition across the Transect

The total number of reads per well varied across the transect, with an average of 21,795 reads per well (range: 3925–66,965 reads, Table 2). The wells sampled for this study contained a total of 2668 unique operational taxonomic units (OTUs), with 4.8% of the total OTUs identified at the species level (128 OTUs out of 2668). On average, there were 250 OTUs detected per collected sample, with 236 different bacterial OTUs and 13 archaeal OTUs identified.

Archaea were observed at a higher average percent abundance than bacteria in most of the 22 sampled wells, ranging from 8.3 to 86.7% of total abundance (Figure 2; Table 2). A methanogen, regardless of the percent abundance of Archaea, was always the dominant OTU in each well, and methanogens, on average, represented 98.6% of the archaean population (Figure S1). Bacteria were generally dominated by Proteobacteria (with Alphaproteobacteria or Gammaproteobacteria found at the highest abundance), Firmicutes, Bacteroidetes, and Actinobacteria (Figure S1). However, only 8 bacterial phyla were observed, on average, at greater than 1% abundance across the transect.

Only 18 different OTUs were present at greater than 10% abundance in any well sampled, 5 of which were methanogens (Figure 3; R commands: as.dendrogram; heatmap). Methanococcus spp. dominated the microbial structure of the 5 shallow, low salinity (up-dip) wells, which were also characterized by low temperatures and high extent of crude oil biodegradation. Methanothermococcus spp. was the most abundant microorganism in 15 of the wells (Figure 3), which were characterized by a wide range of temperatures and chloride concentrations (Table 1). Methanococcus spp. requires much lower growth temperatures than Methanothermococcus spp., possibly (R command: ANOSIM; p = 0.018) explaining the shift in dominance from Methanococcus spp. to Methanothermococcus spp. between the cooler down-dip fields and the warmer up-dip fields (Whitman and Jeanthon, 2006, Figure 4). Methanococccaceae generate methane from either formate or CO₂/H₂ (Liu and Whitman, 2008), confirming previous conclusions that methanogenesis in this region is predominantly hydrogenotrophic (Warwick et al., 2008; McIntosh et al., 2010).

The remaining two wells were dominated by either Methanohalophilus halophilus or Methanolobus spp. The dominance of M. halophilus in well O6 was surprising based on the preferred high salinity growth for the organism—1200 mM Cl⁻ (Kendall and Boone, 2006)—and also because well O6 was not the most saline well sampled. Methanolobus spp., which is present at the highest percent abundance in well NH2, is a methylo trophic methanogen that cannot use H₂/CO₂, acetate, or formate to grow (Kendall and Boone, 2006). Again, no preferential growth conditions for Methanolobus spp. would justify the presence of Methanolobus spp. in well NH2 over other wells across the transect.

Bray-Curtis distances were calculated (R commands: decostand; vegdist) to assess similarity in community composition across the sampled wells (Figure 2). The 5 up-dip, low-salinity, highly biodegraded wells (located in the CR, S, and TU fields) clustered (according to microbial community composition only) away from the remaining 17 wells (R command: hclust), which spanned all ranks of methanogenesis (high, medium, and low). NMDS ordination (also using
| Well ID | Number of Reads* | Species richness* | Pielou's Evenness | % Archaea | % Bacteria | Most Abundant OTU (Archaea) | Most Abundant Bacteria |
|---------|-----------------|-------------------|-------------------|-----------|------------|-----------------------------|-----------------------|
| CR1     | 29,076          | 438               | 0.82              | 59.7      | 40.3       | Methanococcus spp.          | Flexistipes spp.      |
| CR2     | 46,150          | 442               | 0.81              | 75.8      | 24.2       | Methanococcus spp.          | Chromatiales           |
| CR3     | 66,965          | 655               | 0.71              | 52.1      | 47.9       | Methanococcus spp.          | MSBL8                 |
| S3      | 12,964          | 246               | 0.90              | 78.1      | 21.9       | Methanococcus spp.          | Desulfurococcus spp.  |
| TU1     | 35,894          | 316               | 0.79              | 42.5      | 57.5       | Methanococcus spp.          | Clostridiales          |
| O3      | 8557            | 268               | 0.84              | 26.9      | 73.1       | Methanococcus spp.          | Alicyclobacillus spp. |
| O4      | 5929            | 144               | 0.89              | 42.9      | 57.1       | Methanococcus spp.          | Alicyclobacillus spp. |
| O5      | 28,714          | 219               | 0.69              | 65.9      | 34.1       | Methanococcus spp.          | Defibacter spp.       |
| O6      | 45,552          | 138               | 0.82              | 72.2      | 27.8       | Methanohalophilus halophilus | Thermovirgaceae       |
| O7      | 13,711          | 120               | 0.80              | 40.4      | 59.6       | Methanothermococcus spp.    | Acinetobacter lwoffii  |
| O8      | 10,475          | 335               | 0.88              | 38.3      | 61.7       | Methanothermococcus spp.    | Alicyclobacillus spp. |
| O18     | 13,167          | 270               | 0.83              | 53.8      | 46.2       | Methanothermococcus spp.    | Alicyclobacillus spp. |
| O26     | 17,624          | 197               | 0.77              | 55.6      | 44.3       | Methanothermococcus spp.    | Thermovirgaceae        |
| O27     | 39,069          | 166               | 0.60              | 78.3      | 21.7       | Methanothermococcus spp.    | BA201                 |
| O31     | 3925            | 255               | 0.77              | 37.9      | 62.1       | Methanothermococcus spp.    | Halanaerobium spp.     |
| O32     | 5510            | 229               | 0.888             | 8.3       | 91.7       | Methanothermococcus spp.    | Alicyclobacillus spp. |
| NH1     | 4858            | 200               | 0.82              | 29.9      | 70.1       | Methanothermococcus spp.    | Chitinophagaceae       |
| NH2     | 18,430          | 173               | 0.82              | 50.0      | 50.0       | Methanolobus spp.           | Thermovirgaceae        |
| NH3     | 17,603          | 117               | 0.84              | 86.7      | 13.3       | Methanothermococcus spp.    | Alicyclobacillus spp. |
| CB1     | 5966            | 126               | 0.84              | 51.0      | 49.0       | Methanothermococcus spp.    | Methylbacterium kornagate |
| CB3     | 5131            | 166               | 0.87              | 11.0      | 89.0       | Methanothermococcus spp.    | Plesiomonas shigelloides |
| CB4     | 44,233          | 290               | 0.62              | 70.9      | 29.1       | Methanothermococcus spp.    | Calanaerocella spp.    |

*Calculated prior to rarefaction.

†Number of species identified after rarefaction at >0% abundance.

**FIGURE 2** | Cluster dendrogram showing that the 5 up-dip wells (CR, S, and TU) cluster away from the remaining 17 down-dip wells.
Darker colors indicate greater percent abundance than the lighter colors. The taxa found in greatest abundance across the transect are *Methanohalophilus halophilus*, *Methanothermococcus* spp., *Methanococcus* spp., and *Methanolobus* spp. 235 OTUs were identified at >1% abundance in at least one well sampled.

Bray-Curtis distance) also visualized (R command: metaMDS) that the 5 up-dip shallow wells (CR, S, and TU) clustered together. However, they only clustered together when the data were analyzed against 3 of the 11 different environmental parameters, binned chloride concentration, extent of crude oil biodegradation, and oil field/depth (Figure 4), indicating that these parameters may be driving microbial community compositional differences across the transect. The Kruskal-Wallis test (R command: kruskal.test) found that microbial community richness (i.e., number of OTUs present in each sample) was significantly different ($p < 0.05$; Table S2) between groups of samples binned by spatial location within the reservoir (i.e., oil field/depth), pH, and extent of crude oil biodegradation. The number of identified OTUs in the shallow, high pH, highly biodegraded samples was generally greater than those identified in the deeper, lower pH, and low biodegradation wells.

Spatial location within the reservoir, chloride concentrations, and extent of crude oil biodegradation all gave significant $p$-values ($\leq 0.001$), and statistical $R$-values greater than or equal to 0.5 (Table S2) when applying the ANOSIM test (R command: anosim). This means that for those three categorizes, there is a statistical difference between the average community compositions of the binned wells analyzed for each of the categories (i.e., the wells binned by high chloride concentrations have an average microbial community composition that is different than those binned by low chloride concentrations). The ANOSIM test was repeated for just the bacteria identified in each well and just the Archaea identified in each well. When just analyzing the bacteria found in each well, the ANOSIM test only resulted in one significant grouping—by chloride concentration (Table S2). When just analyzing the archaea, the ANOSIM test identified two different significant groupings—extent of crude oil biodegradation and alkalinity. This suggests that both the Archaea and bacteria found in each well may be driving the amount of crude oil biodegradation found in each well, that crude oil biodegradation was accompanied by microbial community growth, and/or that the Archaea may impact alkalinity concentrations, independent of the bacteria.
Microbial Community Differences between the up-dip and down-dip Samples

Due to the sample clustering visualized using NMDS (e.g., the 5 up-dip wells clustered away from the 17 other wells), these 5 samples were removed from the statistical analysis. The ANOSIM test was performed again to determine if those 5 up-dip samples were driving the microbial community differences observed across the transect, or if there were still microbial community structure differences in the remaining 17 wells (i.e., is the variability identified exclusive to the 5 up-dip samples?). When removing those 5 samples, hydrogeochemical differences were still abundant across the remaining 17 samples; the remaining 17 wells span 3 different oil fields, all extents of methanogenesis (high, medium, and low), produced fluids with temperature values from 24.4 to 48.5°C, pH values between 6.7 and 7.4, δ^{13}C-DIC values from +0.2 to +24.1‰, alkalinity values between 4.7 and 56.4 meq/kg, and chloride concentrations between 1154 and 2056 mM (Table 1). After removing those 5 samples and re-running the ANOSIM test, no environmental parameters were associated with significantly different microbial structures.
(p > 0.05; Table S2). This indicates that the major microbial community variations are found in the 5 up-dip samples, and that the remaining 17 samples have statistically similar microbial communities when binned by environmental parameters.

When analyzing the average order-level taxonomy of the two groups of samples (5 up-dip wells vs. the 17 down-dip wells), there were some major differences in microbial structure. Thirty-five different orders were identified at, on average, greater than 0.5% abundance in either of the two groups of samples. *Methanococcales* was the dominant order in both groups of samples; however, *Methanococcus* spp. dominated the up-dip wells while *Methanothermococcus* spp. dominated the down-dip wells (Section Differences in Microbial Community Composition across the Transect). No other identified orders were found at greater than 10% abundance in both groups. Some specific orders were found at a much higher percent abundance (or are only present) in the up-dip wells, and vice-versa. Most notably, greater than 90% of the identified sequences of *Cloacamonales*, *Bacteroidales*, *Chromatiales*, *Desulfobacterales*, *Desulfuromonadales*, *Methanobacterales*, OPB11, PL-11B10, *Spirochaetales*, and YLA114 were found in the up-dip oil fields, while greater than 90% of the identified sequences of *Halanaerobiales*, *Oceanaeruphylaceae*, and *Synergistales* were found in the down-dip wells.

All of the orders containing sulfate reducing bacteria (SRB) were found in much higher abundance in the up-dip, low salinity, highly biodegraded, non-methanogenic wells. These include *Desulfobacterales*, *Desulfovirionales*, *Desulfuromonadales*, *Methanobacterales*, OPB11, PL-11B10, *Spirochaetales*, and YLA114 were found in the up-dip oil fields, while greater than 90% of the identified sequences of *Halanaerobiales*, *Oceanaeruphylaceae*, and *Synergistales* were found in the down-dip wells.

Factors Driving Microbial and Metabolic Differences across the Transect

These results indicate that the extent of methanogenesis, which varies widely across the sampled transect, is not impacted by either the bacterial or archaeal communities present in each well. This may suggest that although “who’s there” (i.e., the microbial communities identified at the time of sampling) may not have influenced the amount of methanogenesis, that other factors (possibly salinity or actual abundances of archaea and bacteria) may control whether or not certain communities are active and will generate methane. It is also possible that the microbial community has changed dramatically over time, and that the microbes present during the onset of crude oil biodegradation and methanogenesis may not have been captured by our sampling efforts. However, the extent of crude oil biodegradation may indeed be impacted by the methanogens present, as
indicated by the results of the ANOSIM test. This evidence would further confirm that crude oil biodegradation and methanogenesis are linked in subsurface petroleum reservoirs (e.g., Jones et al., 2008; Gieg et al., 2010; Shelton et al., 2016), and that Archaea may play a major role in degrading the hydrocarbons, not just in generating methane.

Current research also suggests that there may not be major differences in the microbial communities present in biodegraded vs. non-biodegraded reservoirs (Sette et al., 2007; Oliveira et al., 2008), and that the major difference may be the relative abundance of the populations present (i.e., species evenness). This conclusion is not in agreement with the findings from this study. Bacterial richness is much higher on average in the oil fields exhibiting higher amounts of crude oil biodegradation (up-dip fields) than ones exhibiting minimal crude oil alteration (down-dip fields). The average Pielou evenness value (Table 2) is also similar between the 5 up-dip (0.81) and the 17 down-dip (0.80) samples, indicating similar evenness for the highly biodegraded samples compared to the less biodegraded samples. Furthermore, there are significantly different microbial community structures between the highly biodegraded and moderately/minimally biodegraded areas of the reservoir, indicating different microbial communities (and differences in their abundance) surrounding these wells. However, this could be due to the hydrogeochemistry of the reservoir, as the more degraded fields are associated with lower temperature and more dilute fluids than the deeper oil fields.

Salinity has been shown to be a major driver of microbial community presence and metabolism, affecting both crude oil biodegradation and microbial methanogenesis (e.g., Potter et al., 2009; Oren, 2011; Head et al., 2014). Shelton et al. (2016) suggested that chloride concentrations were driving differences in the amount of crude oil biodegradation across the transect sampled in this study. This study also provides evidence to support salinity as a driver of microbial community differences (as shown by a significant ANOSIM p-value). There are two major changes in salinity concentrations across the transect: between the shallow wells and the mid-depth wells, and between the mid-depth well and the deep wells.

However, it is important to note that major changes in salinity, spatial location, and extent of crude oil biodegradation exist when comparing the 17 down-dip wells, even though there appears to be no major microbial structure differences across these 17 wells. The 17 down-dip wells span three different oil fields and 921 m of depth, produce formation water with chloride concentrations between 1154 and 2056 mM (Table 1), and produce crude oil that is either moderately or minimally biodegraded. Therefore, it is difficult to say that, “salinity is driving microbial community differences across the transect,” because there are major salinity differences present down-dip, with fairly similar average microbial communities present along that gradient.

It is more prudent to hypothesize that a salinity threshold may exist between 776 and 1154 mM that impacted the microbial community structure in this formation. Crude oil biodegradation may be limited above this hypothesized threshold, because there is a major difference in crude oil biodegradation between wells producing water with these different chloride concentrations. It is difficult to determine if methanogenesis itself is limited at lower salinity values, or if methanogens have been (or are being) outcompeted by SRB around these wells locations. Therefore, microbial function in these 5 up-dip wells may not be limited by salinity specifically, but rather by the lack of sulfate present to support the SRB identified in the 5 up-dip wells.

High concentrations of salinity may not impact microbial community structure, as indicated by the results of this study (e.g., the 17 down-dip wells), but merely microbial function (i.e., metabolism and roles). Both crude oil biodegradation and microbial methanogenesis are limited in the most-saline field sampled, the field with wells producing formation water with chloride concentrations >2000 mM. As the microbial community structure in the highly-saline fields is statistically similar to the structure of the centrally located methanogenic field, it is fair to assume that salinity is limiting the lack of function in these down-dip fields (i.e., limiting methanogenesis and/or crude oil biodegradation).

CONCLUSIONS

Archaea, on average, dominated the microbial community composition in each sampled location, while a methanogen, regardless of the percent Archaea, was the dominating microbial species in each sampled well location. Statistical analyses performed on the total community structure suggested that chloride concentrations, spatial location within the Gulf Coast Basin (i.e., well depth and surficial location), and the extent of biodegradation have the most influence on the microbial community structure across the transect. However, salinity and spatial location within the reservoir are co-linear, and are associated with one another. The statistical variability between microbial structure and environmental parameters is due mostly to processes occurring in 5 specific well locations, where the shallowest wells produce fluids with the lowest chloride concentrations and highest extent of crude oil biodegradation. After removing the data from these 5 wells from the statistical analyses, no significant differences in microbial community structure exist when the remaining 17 wells are binned by environmental parameters, even though environmental gradients still exist (i.e., changes in salinity, temperature, and pH are prevalent). Therefore, we hypothesize that either (1) a chloride threshold (between approximately 800–1000 mM) exists, resulting in a dramatic shift in the microbial community structure between the 5 up-dip well locations and the 17 down-dip well locations; and/or (2) the major difference between these two groups of sample locations is due to differences in crude oil biodegradation, likely stemming from variations in metabolism (sulfate reduction vs. methanogenesis). Furthermore, this salinity threshold may also be driving the high amounts of crude oil biodegradation observed in the low salinity, up-dip well locations, leading to the conclusion that salinity is the major driver in controlling microbiology across the transect. Salinity does not appear to impact microbial community structure beyond this threshold, as indicated by the lack of major
difference in microbial community structure in the down-dip wells. Variations in methanogenesis across the transect are not linked to statistically different microbial communities, and therefore, as hypothesized in Shelton et al. (2016), are likely due to hydrogeochemistry and oil quality.

This research has implications for enhancement of natural gas production from crude oil biodegradation. Although, methanogens were abundant, represented by low species diversity, and found in every well sampled for this study regardless of the apparent extent of methanogenesis, extent of methanogenesis still varies across the transect. This implies that engineering reservoirs to favor specific methanogen populations may not be useful when trying to accelerate methanogenic crude oil biodegradation, because hydrogeochemical conditions of the reservoir may be affecting the methanogens’ ability to convert organic substrates to natural gas. Crude oil biodegradation is likely limited by either salinity or by microbial ecology, as both change drastically between the 5 up-dip, low-salinity, highly biodegraded samples and the remaining 17 down-dip, higher salinity, low biodegradation samples. Methanogenesis seems to be more halotolerant than crude oil biodegradation, however the hypothesized switch from sulfate reduction to methanogenesis in the shallow, up-dip fields would need to be further investigated to determine if methanogenesis will occur in this low-salinity, low oil quality environment.

AUTHOR CONTRIBUTIONS

JLS helped devise the research plan, performed field work, analyzed samples and resulting data, and wrote and edited the manuscript. DA assisted in devising the research plan, analyzing the samples and resulting data, and edited the manuscript. JCM assisted in devising the research plan, analyzing the data, and edited the manuscript. JRS assisted in devising the research plan, analyzing the data, and in editing the manuscript. NF assisted in devising the research plan and in editing the manuscript. JEM assisted in devising the research plan, analyzing the data, and in editing the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2016.01535

REFERENCES

Berdugo-Clavijo, C., Dong, X., Soh, J., Sensen, C. W., and Gieg, L. M. (2012). Methanogenic biodegradation of two-ringed polycyclic aromatic hydrocarbons. FEMS Microbiol. Ecol. 81, 124–133. doi: 10.1111/j.1574-6941.2012.01328.x

Berdugo-Clavijo, C., and Gieg, L. M. (2014). Conversion of crude oil to methane by a microbial consortium enriched from oil reservoir production waters. Front. Microbiol. 5:197. doi: 10.3389/fmicb.2014.00197

Bethke, C. M., Sanford, R. A., Kirk, M. F., Jin, Q., and Flynn, T. M. (2011). The thermodynamic ladder in geomicrobiology. Am. J. Sci. 311, 183–210. doi: 10.2475/03.2011.01

Bonin, A. S., and Boone, D. R. (2006). “The Order Methanobacteriales,” in The Prokaryotes, eds M. Dworkin, S. Falkow, E. Rosenberg, K. H. Schleifer, and E. Stackebrandt (New York, NY: Springer), 231–243.

Carr, S. A., Orcutt, B. N., Mandernack, K. W., and Spear, J. R. (2015). Abundant Atribacteria in deep marine sediment from the Adélie Basin, Antarctica. Front. Microbiol. 6:872. doi: 10.3389/fmicb.2015.00872

Dahle, H., Garshol, F., Madsen, M., and Birkeland, N. K. (2008). Microbial community structure analysis of produced water from a high-temperature North Sea oil-field. A. Van Leeuwen. 93, 37–49. doi: 10.1007/s10482-007-9177-z

Edgar, R. C. (2013). UPARSE: highly accurate OTU sequences from microbial amplicon reads. Nature Meth. 10, 996–998. doi: 10.1038/nmeth.2604

Fierer, N., Leff, J. W., Adams, B. J., Nielsen, U. N., Bates, S. T., Lauber, C. L., et al. (2012). Cross-biome metagenomic analyses of soil microbial communities and their functional attributes. Proc. Natl. Acad. Sci. U.S.A. 109, 21390–21395. doi: 10.1073/pnas.1215210110

Gieg, L. M., Davidova, I. A., Duncan, K. E., and Sulflita, J. M. (2010). Methanogenesis, sulfate reduction and crude oil biodegradation in hot Alaskan oilfields. Env. Micro. 12, 3074–3086. doi: 10.1111/j.1462-2920.2010.02282.x

Gieg, L. M., Duncan, K. E., and Sulflita, J. M. (2008). Bioenergy production via microbial conversion of residual oil to natural gas. Appl. Environ. Microbiol. 74, 3022–3029. doi: 10.1128/aem.00119-08

Gieskes, J. M., and Rogers, W. C. (1973). Alkalinity determination in interstitial waters of marine sediments. J. Sediment. Res. 43, 272–277.

Göker, M., Teshima, H., Lapidus, A., Nolan, M., Lucas, S., Hammon, N., et al. (2011). Complete genome sequence of the acetate-degrading sulfate reducer Desulfofusca acetoxidans type strain (ASRB2T). Stand. Genomic Sci. 4, 393–401. doi: 10.4056/sigs.206705

Grabowski, A., Nercissian, O., Fayolle, F., Blanchet, D., and Jeanthon, C. (2005). Microbial diversity in production waters of a low-temperature biodegraded oil reservoir. FEMS Microbiol. Ecol. 54, 427–443. doi: 10.1016/j.femsec.2005.03.009

Head, I. M., Gray, N. D., and Larter, S. R. (2014). Life in the slow lane; biogeochemistry of biodegraded petroleum containing reservoirs and implications for energy recovery and carbon management. Front. Microbiol. 5:566. doi: 10.3389/fmicb.2014.00566

Hein, F. J., Leckie, D., Larter, S., and Suter, J. R. (2013). “Heavy oil and bitumen petroleum systems in Alberta and beyond: the future is nonconventional and the future is now,” in Heavy- Oil and Oil-Sand Petroleum Systems in Alberta and Beyond: AAPG Studies in Geology, Vol. 64, eds F. J. Hein, D. Leckie, S. Larter, and J. R. Suter (Tulsa, OK: American Association of Petroleum Geologists), 1–21.

Hu, P., Tom, L., Singh, A., Thomas, B. C., Baker, B. J., Piceno, Y. M., et al. (2016). Genome-resolved metagenomic analysis reveals roles for candidate phyla and
other microbial community members in biogeochemical transformations in oil reservoirs. *mBio* 7:e01669-15. doi: 10.1128/mBio.01669-15

Hug, L. A., Castelle, C. J., Wrighton, K. C., Thomas, B. C., Sharon, I., Frischkorn, K. R., et al. (2013). Community genomic analyses constrain the distribution of metabolic traits across the Chloroflexi phylum and indicate roles in sediment carbon cycling. *Microbiome* 1:22. doi: 10.1186/2049-2618-1-22

Jones, D. M., Head, I. M., Gray, N. D., Adams, J. J., Rowan, A. K., Aitken, C. M., et al. (2008). Crude-oil biodegradation via methanogenesis in subsurface petroleum reservoirs. *Nature* 451, 176–180. doi: 10.1038/nature06484

Kendall, M. M., and Boone, D. R. (2006). “The order Methanosarcinales,” in *The Prokaryotes*, eds M. Dworkin, S. Falkow, E. Rosenberg, and K. H. Evans (New York, NY: Springer), 244–256.

Kirk, M. F., Wilson, B. H., Marquart, K. A., Zeglin, L. H., Vinson, D. S., and Flynn, T. M. (2015). Solute concentrations influence microbial methanogenesis in coal-bearing strata of the Cherokee basin, USA. *Front. Microbiol.* 6:1287. doi: 10.3389/fmicb.2015.01287

Kleinsteuber, S., Schleinitz, K. M., and Vogt, C. (2012). Key players and team play: anaerobic microbial communities in hydrocarbon-contaminated aquifers. *App. Microbiol. Biotech.* 94, 851–873. doi: 10.1007/s00253-012-4025-0

Kryachko, Y., Dong, X., Sensen, C. V., and Vooroud, G. (2012). Compositions of microbial communities associated with oil and water in a mesothermal oil field. *A. Van Leeuwen. J. Microb*. 101, 493–506. doi: 10.1007/s10482-011-9568-y

Kuever, J. (2014). “The family Syntrophorhabdaceae,” in *The Prokaryotes*, ed E. Rosenberg (Berlin: Heidelberg: Springer), 301–303.

Larter, S., Huang, H., Adams, J., Bennett, B., Jokanola, O., Oldenburg, T., et al. (2006). The controls on the composition of biodegraded oils in the deep subsurface: Part II—Geological controls on subsurface biodegradation fluxes and constraints on reservoir-fluid property prediction. *AAPG Bull.* 90, 921–938. doi: 10.1306/01270605130

Larter, S., Wilhelms, A., Head, I., Koopmans, M., Aplin, A., Di Primio, R., et al. (2003). The controls on the composition of biodegraded oils in the deep subsurface—part 1: biodegradation rates in petroleum reservoirs. *Org. Geochem.* 34, 601–613. doi: 10.1016/S0146-6380(02)00240-1

Li, W., Wang, L. Y., Duan, R. Y., Liu, J. F., Gu, J. D., and Mu, B. Z. (2012). Microbial community characteristics of petroleum reservoir production water amended with an alkane and incubated under nitrate-, sulfate-reducing and methanogenic conditions. *Int. Biodeterior. Biodegr.* 69, 87–96. doi: 10.1016/j.ibiod.2012.01.005

Limam, R. D., Chouari, R., Mazéas, L., Wu, T. D., Li, T., Grossin-Debatista, J., et al. (2014). Members of the uncultured bacterial candidate division WWE1 are implicated in anaerobic digestion of cellulose. *Microb. Open* 3, 157–167. doi: 10.1002/mbo3.144

Liu, Y., and Whitman, W. B. (2008). Metabolic, phylogenetic, and ecological diversity of the methanogenic archaea. *Ann. N.Y. Acad. Sci.* 1125, 171–189. doi: 10.1196/annals.1419.019

Mayumi, D., Dolfing, J., Sakata, S., Maeda, H., Miyagawa, Y., Ikarashi, M., et al. (2013). Carbon dioxide concentration dictates alternative methanogenic pathways in oil reservoirs. *Nat. Commun.* 4, 1998. doi: 10.1038/ncomms2998

McDonald, D., Clemente, J. C., Kuczynski, J., Rideout, J. R., Stombaugh, J., Wendel, D., et al. (2012). The Biological Observation Matrix (BIOM) format or: how I learned to stop worrying and love the ome-ome. *GigaScience* 1:7. doi: 10.1186/2047-217X-1-7

McIntosh, J. C., Warwick, P. D., and McCray, J. M. (2016). Impacts of formation water geochemistry and crude oil biodegradation on microbial methanogenesis. *Org. Geochem.* 98, 105–117. doi: 10.1016/j.orggeochem.2016.05.008

Shelton, J. L., McIntosh, J. C., Warwick, P. D., and Yi, A. L. Z. (2014). Fate of injected CO2 in the Wilcox Group, Louisiana, Gulf Coast Basin: chemical and isotopic tracers of microbial brine–rock CO2 interactions. *App. Geochem.* 51, 155–169. doi: 10.1016/j.apgeochem.2014.09.015

Wang, L. Y., Gao, C. X., Mbadinga, S. M., Zhou, L., Liu, J. F., Gu, J. D., et al. (2011). Characterization of an alkane-degrading methanogenic enrichment culture from production water of an oil reservoir after 274 days of incubation. *Int. Biodeter. Biodegr.* 65, 444–450. doi: 10.1016/j.ibiod.2010.12.010

Whitman, P. D., Breland, F. C., and Hackley, P. C. (2008). Biogenic origin of coalbed gas in the northern Gulf of Mexico Coastal Plain, USA. *Int. J. Coal Geol.* 76, 1–2. doi: 10.1016/j.coal.2008.05.009

Whitman, W. B., Bowen, T. L., and Boone, D. R. (2006). “The methanogenic bacteria,” in *The Prokaryotes*, ed M. Dworkin (New York, NY: Springer), 165–207.

Whitman, W. B., and Jeantonn, C. (2006). “Methanococcales,” in *The Prokaryotes*, ed M. Dworkin (New York, NY: Springer), 257–273.

Yamane, K., Hattori, Y., Ohtagaki, H., and Fujisawa, K. (2011). Microbial diversity with dominance of 16S rRNA gene sequences with high GC contents at 74 and 98 C-subsurface crude oil deposits in Japan. *JEMS Microbiol. Ecol.* 76, 220–235. doi: 10.1111/j.1365-7134.2011.01044.x

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