

ORIGINAL ARTICLE

Biostimulation induces syntrophic interactions that impact C, S and N cycling in a sediment microbial community

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Stimulation of subsurface microorganisms to induce reductive immobilization of metals is a promising approach for bioremediation, yet the overall microbial community response is typically poorly understood. Here we used proteogenomics to test the hypothesis that excess input of acetate activates complex community functioning and syntrophic interactions among autotrophs and heterotrophs. A flow-through sediment column was incubated in a groundwater well of an acetate-amended aquifer and recovered during microbial sulfate reduction. *De novo* reconstruction of community sequences yielded near-complete genomes of *Desulfobacter* (*Deltaproteobacteria*), *Sulfurovum*- and *Sulfurimonas*-like *Epsilonproteobacteria* and *Bacteroidetes*. Partial genomes were obtained for *Clostridiales* (*Firmicutes*) and *Desulfuromonadales*-like *Deltaproteobacteria*. The majority of proteins identified by mass spectrometry corresponded to *Desulfobacter*-like species, and demonstrate the role of this organism in sulfate reduction (Dsr and APS), nitrogen fixation and acetate oxidation to CO₂ during amendment. Results indicate less abundant *Desulfuromonadales*, and possibly *Bacteroidetes*, also actively contributed to CO₂ production via the tricarboxylic acid (TCA) cycle. Proteomic data indicate that sulfide was partially re-oxidized by *Epsilonproteobacteria* through nitrate-dependent sulfide oxidation (using Nap, Nir, Nos, SQR and Sox), with CO₂ fixed using the reverse TCA cycle. We infer that high acetate concentrations, aimed at stimulating anaerobic heterotrophy, led to the co-enrichment of, and carbon fixation in *Epsilonproteobacteria*. Results give an insight into ecosystem behavior following addition of simple organic carbon to the subsurface, and demonstrate a range of biological processes and community interactions were stimulated.

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Introduction

The role of microorganisms in organic carbon cycling is of considerable interest, as these activities can impact carbon turnover rates and sequestration (Karl *et al.*, 2012) and influence the fate of contaminants, such as petroleum (Hazen *et al.*,

2010), uranium (Anderson *et al.*, 2003) or arsenic (Islam *et al.*, 2004). Sediments host a substantial proportion of the environmental bacterial and archaeal biomass, with an estimated 6–40% of the prokaryotic biomass inhabiting the terrestrial subsurface, and an estimated total carbon content rivaling that of the global plant biomass (Whitman *et al.*, 1998). Most endogenous organic matter in terrestrial sediment is considered to be relatively refractory, consisting largely of polymers, such as lignin and cellulose, and plant-derived (or even petroleum-based) hydrocarbons (Hartog *et al.*, 2004; Rowland *et al.*, 2006). However, some fractions of complex organic matter can be degraded by fermentative or respiratory microorganisms (Benner *et al.*, 1984; Leschine, 1995; Widdel and Rabus, 2001).

In anaerobic environments, such as the terrestrial subsurface, acetate is an important product of central carbohydrate degradation pathways,

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following the oxidation of pyruvate. It is also a common product of respiration or fermentation of other organic acids and alcohols by anaerobic microorganisms, including sulfate-reducing bacteria (SRB) that incompletely oxidize organic matter (Gibson, 1990). Characteristic pathways identified through which acetate oxidation to CO₂ proceeds are the TCA cycle and the acetyl-CoA pathway (Thauer *et al.*, 1989). Both pathways are reversible and can be used by autotrophic bacteria to catalyze the reduction of CO₂ to back to acetyl-CoA (Fuchs, 1986). Acetate can be used in methanogenesis, and numerous bacteria can couple acetate oxidation to the reduction of inorganic compounds, such as Fe(III), U(VI), nitrate, S⁰ and sulfate (Thauer *et al.*, 1989; Lovley *et al.*, 2004). As a biologically relevant and non-fermentable organic compound, acetate is an attractive substrate to use in studying biogeochemical cycling linked to anaerobic respiration and bioremediation (Anderson *et al.*, 2003).

While relatively few organisms proliferate when stimulating sediment microbial communities with acetate (Holmes *et al.*, 2002), the enriched organisms constitute a community that is nonetheless intricate (Handley *et al.*, 2012). Community processes can be expected to include syntrophic interactions, biomass recycling, and the precipitation of diverse and in some cases, cryptic biogeochemical cycles. Unraveling the individual roles of community members, and their metabolic pathways is non-trivial.

Here, we used genomics-informed proteomic information (proteogenomics, Ram *et al.*, 2005) to resolve organism-specific activity and detect metabolic pathways utilized in an acetate-amended, sediment-hosted subsurface microbial community, *a priori*. The study provides a snapshot of community-wide functioning and interactions in an aquifer setting during acetate-induced uranium bioremediation under predominantly sulfate-reducing conditions, and builds on previous studies considering the role of bacteria, in particular *Geobacter*, in Fe(III) and U(VI) reduction within the aquifer (for example, Anderson *et al.*, 2003; Holmes *et al.*, 2009; Wilkins *et al.*, 2009; Williams *et al.*, 2011). Our proteogenomic data demonstrate the activity of bacteria linked to metal reduction, in addition to microbial utilization of the carbon, nitrogen and sulfur, and the indirect stimulation of autotrophic (or mixotrophic) bacteria in response to CO₂ and sulfide generated through acetate-dependent sulfate reduction. Biogeochemical reaction modeling was also used to evaluate denitrification pathways.

Materials and methods

Experiment setup and sampling

Un-amended and acetate-amended sediments were sampled from an alluvial freshwater aquifer underlying the Department of Energy's Integrated Field

Research Challenge (IFRC) site at Rifle, CO, USA. In order to stimulate aquifer sediment with acetate *in situ*, and access sediment from the subsurface post-stimulation, we incubated sediment in a flow-through column in an existing groundwater well (P104; see well gallery in Williams *et al.*, 2011). Un-amended sediment was first excavated from the aquifer using a backhoe, and sieved (to remove rocks) to a final particle size of <2 mm. This material was packed into a clear custom built PVC cylindrical column (5.1 cm wide × 10.2 cm long) and incubated ~5 m below ground surface within the well (Supplementary Figure S1). The column equilibrated with subsurface conditions for 15 days before amendment, which have previously been shown to be anoxic with <16 μM of dissolved oxygen (Williams *et al.*, 2011). Acetate (electron donor and carbon source) and bromide (conservative tracer) were injected into wells ~0.5 m upgradient of well P104, obtaining final concentrations of ~15 and ~1.3 mM, respectively (method in Williams *et al.*, 2011). Amended groundwater was pumped up through the column for 24 days, as described previously (Handley *et al.* 2012), during which this region of the aquifer was subject to its third consecutive summer of acetate amendment. Sediment from the entire column was homogenized and then flash frozen upon collection. Comparative analyses with a replicate column and other aquifer samples are described elsewhere (Handley *et al.*, 2012).

Geochemistry analyses

Groundwater samples were filtered using 0.25 μm PTFE filters for geochemical analyses. Acetate, bromide, sulfate, chloride, uranium, Fe(II) and sulfide were measured, as described previously (Williams *et al.* 2011).

DNA extraction

Genomic DNA was extracted from 5 g of un-amended sediment and 300 g of acetate-amended sediment (5–10 g per tube) using PowerMax Soil DNA Isolation Kits (MoBio Laboratories, Inc., Carlsbad, CA, USA) with the following modification to the manufacturer's instructions. Sediment was vortexed at maximum speed for an additional 2 min in SDS, and then incubated for 30 min at 60 °C instead of bead beating. Acetate-amended sediment extraction replicates were pooled in order to obtain ~8 μg of DNA for downstream analysis. All eluted DNA was concentrated, as described by Handley *et al.* (2012).

16S rRNA gene clone libraries

DNA was amplified using the general bacterial 16S rRNA gene primers 27f and 1492r (Lane, 1991), and a temperature gradient to minimize PCR bias, which comprised 11 PCR reactions at 11 different annealing temperatures. The PCR protocol was: 1 min at

94 °C; 25 cycles of 1 min at 94 °C, 30 s at 48–58 °C (11 temperature gradient) and 1 min at 72 °C; and 7 min at 72 °C. Amplicons were pooled, and precipitated as above. Clone libraries were constructed using the TOPO TA Cloning Kit (Invitrogen, Carlsbad, CA, USA) with electrocompetent cells. 16S rRNA genes from transformed colonies were PCR amplified using the protocol: 10 min at 95 °C; 25 cycles of 30 s at 95 °C, 30 s at 53 °C and 1.5 min at 72 °C; and 7 min at 72 °C. Inserts were screened for correct size (~1400 bp) by gel electrophoresis, and sequenced using capillary electrophoresis (Applied Biosystems 3730xl DNA Analyzer, Foster City, CA, USA), with M13f (–21) and M13r (–24) primers (Invitrogen). Sequences were trimmed to remove Phred quality scores ≤ 20 , and forward and reverse strands were merged into near full-length sequences using Phrap (<http://www.phrap.org/phredphrapconsed.html>). USEARCH (Edgar, 2010) was used to check for chimeras—after which 98 sequences were retained for each sample (un-amended and acetate-amended)—and to cluster sequences into OTUs 97% similar. A representative from each OTU was BLASTed (Altschul *et al.*, 1990) to a non-redundant version of SILVA SSURef102 (<http://www.arb-silva.de/>).

Metagenomic 16S rRNA gene sequence reconstruction

16S rRNA gene fragments from Illumina metagenomic sequencing (described below) were reconstructed into near full-length genes using EMIRGE (Miller *et al.*, 2011) with 120 iterations and the non-redundant SILVA SSURef102 as the starting database. In the final iteration, 12 032 paired reads (0.05%) were reconstructed into 16S rRNA gene sequences. Sequences were clustered into OTUs $\geq 97\%$ similar. To exclude less reliable rare sequences, OTUs with raw relative abundances $\geq 0.5\%$ were used. Representative sequences were BLASTed to the SILVA database. OTU abundances, calculated on the basis of a probabilistic accounting of read depth, were normalized by sequence lengths.

Phylogenetic analyses

For 16S rRNA phylogenetic analysis genes were aligned to reference sequences from GenBank with Clustal W (Thompson *et al.*, 1994), and Maximum Likelihood trees were created (see Handley *et al.*, 2012). Bootstrap consensus trees were inferred from 1000 replicates. Branches corresponding to partitions reproduced in $< 50\%$ bootstrap replicates were collapsed. Trees were annotated with data sets in iTOL (Letunic and Bork, 2006).

Metagenome sequencing and assembly

Genomic DNA from the acetate-amended sediment was sequenced on one flow-cell of an Illumina Genome Analyzer IIx (Illumina, Inc., San Diego, CA, USA). A paired-end (PE) shotgun library with 1-kb insert size was prepared using Illumina's Genomic

DNA Sample Prep kit, according to the manufacturer's instructions. Sequencing produced ~7 Gb of sequence (*versus* ~400 Mb with 454), and 29 million paired-end reads ~125-bp long. See Supplementary Information for further details, and a description of 454 sequencing and assembly. Illumina reads were trimmed to remove low-quality bases from the 3' ends, after which 87% of paired/single reads > 60 -bp long were retained. Reads were initially assembled eight times using Velvet 1.1 (Zerbino and Birney, 2008) with different parameters optimized on the basis of expected genome coverage, after which a reference-guided Velvet-Columbus re-assembly was undertaken using paired-end scaffolds from the most abundant organism (r9c1, see binning section below and Supplementary Information for details).

Genome annotation

Genes were predicted, and translated into protein sequences using Prodigal (Hyatt *et al.*, 2010), and annotated using the pipeline, as described by Yelton *et al.* (2011).

Genomic binning of metagenomic assemblies

Emergent self-organizing maps of tetranucleotide frequencies and read coverage data were used to define bins (Dick *et al.*, 2009). Sequence fragments 5-kb long were used to create the map and separate sequences into bins, after which 2-kb fragments were projected onto the map. Emergent self-organizing maps (ESOM) were created using Databionics ESOM Tools (Ultsch and Mörchen, 2005). PE-scaffolds > 2 -kb long were also evaluated on the basis of read coverage (that is, relative abundance), gene GC-content and BLASTP best matches to the Uniref90 database (Suzek *et al.*, 2007) to help resolve unclear ESOM bin boundaries and classify low-abundance organisms. EMIRGE-reconstructed 16S rRNA sequences were assigned to genomic bins on the basis of the nearest BLAST-determined database match, read coverage and sequence abundance.

Genome completeness

Genome completeness was determined on the basis of 35 single-copy orthologous groups (OGs) (Raes *et al.*, 2007). OGs were obtained from eggNOG v. 3.0, a database of OGs in 1133 taxonomically diverse organisms, including 943 Bacteria (Powell *et al.*, 2012). Sequences were considered orthologous on the basis of reciprocal BLASTP analysis, if they had a minimum bit score of 60, an alignment length of $\geq 70\%$, and $\geq 30\%$ shared identity.

Whole-genome comparisons

Relationships of assembled genomes to closely related genomes were measured using the amino-acid percent identity averaged across putative

orthologs, determined using reciprocal BLASTP with the same criteria as for genome completeness estimates.

Proteomics

Proteins were extracted from ~10 g of un-amended and acetate-amended sediment *via* a heat-assisted SDS-based method, followed by TCA protein precipitation/acetone washes, trypsin proteolysis, desalting and solvent exchange (Chourey *et al.*, 2010). Digested peptides were loaded in triplicate onto 5 cm strong cation-exchange columns, and connected to a reverse-phase (C18) front column (Phenomenex, Torrance, CA, USA) with an integrated nanospray tip (New Objective, Inc., Woburn, MA, USA). Proteomes were analyzed *via* two-dimensional 24-hour separations with $\text{CH}_3\text{COONH}_4$ salt pulses followed by reverse-phase gradients (Dionex U3000 HPLC, Sunnyvale, CA, USA) with online electrospray tandem mass spectrometry (2D-LC-MS/MS, LTQ Velos Orbitrap, Thermo Scientific, San Jose, CA, USA). Data-dependent MS/MS spectra were acquired, with full scans (400–1700 m/z) at 30-K resolution (top-ten method; dynamic exclusion: one; see VerBerkmoes *et al.*, 2009). MS/MS spectra were queried with SEQUEST (Eng *et al.*, 1994), against a predicted protein database constructed from the Velvet-Columbus metagenome assembly, along with common contaminants. Identified peptide sequences were reassembled into proteins and filtered with DTASelect (Tabb *et al.*, 2002) using the following parameters: Xcorr values >1.8 (+1), 2.5 (+2) and 3.5 (+3) and DeltCN values >0.08, and a requirement for ≥ 2 peptides per locus. The DTASelect and metagenomic database files are available at https://compbio.ornl.gov/ersp_rifle/column_sediment_2009. Summed triplicate technical replicates and normalized spectral abundance factors (NSAFs, Florens *et al.*, 2006) were used for subsequent analyses.

Modeling

The Supplementary Information describes a combined thermodynamic–kinetic modeling approach for microbially mediated rate laws on the basis of work of Jin and Bethke (2005) and Dale *et al* (2008).

Simulations of microbially mediated biogeochemical reactions (see below) were on the basis of physical and acetate-delivery parameters for well P104 and site geochemical data (Figure 1). The system was treated as one dimensional, with average flow and acetate delivery rates estimated from the arrival time of the bromide tracer (3.37 cm per day). Nitrate is present in micromolar concentrations within the aquifer near well P104 (Williams *et al.*, 2011), and varies 2 orders of magnitude across the site (unpublished data). For modeling, we used upper and lower values of $72 \mu\text{M}$ and $5 \mu\text{M}$ nitrate,

with the latter value being more representative of the aquifer local to P104.

The rate of denitrification is affected by the maximum rate per unit cell or mass of biomass, v_{max} , and the biomass concentration, B_{mass} . Lacking cell-count information for the sulfur-oxidizing bacteria, we assumed a rate constant that implicitly accounts for the biomass and results in substantial denitrification over the 0.5-m flow path. We assumed the autotrophic (sulfide-dependent) denitrification reaction rate to be five times slower than heterotrophic denitrification, following Koenig and Liu (2001 and references therein) and Cardoso *et al.*

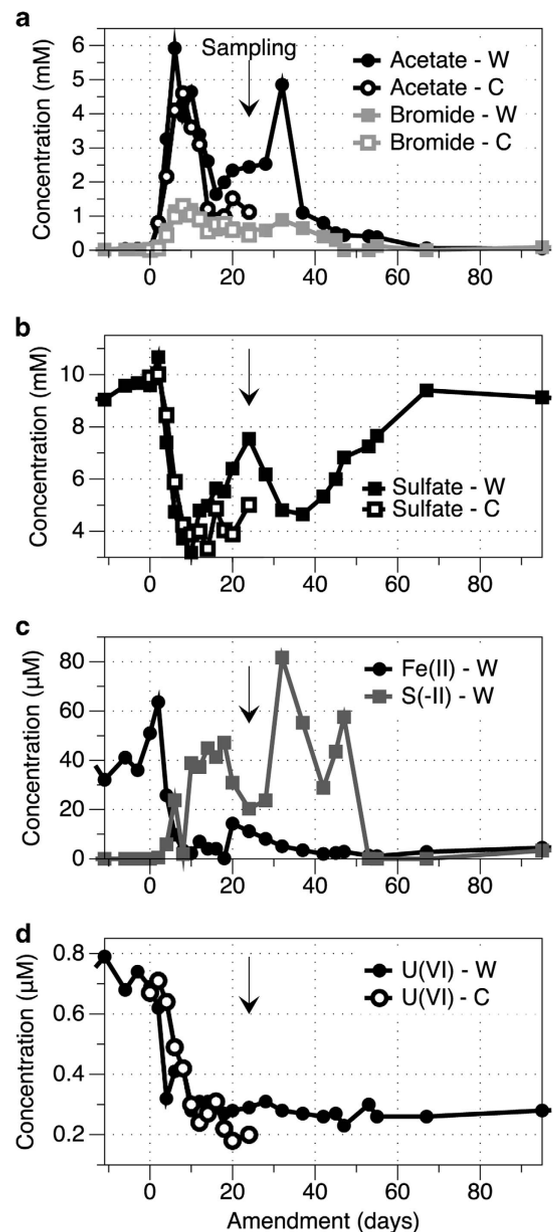


Figure 1 Plot showing concentrations of (a) acetate and bromide, (b) sulfate, (c) Fe(II) and sulfide, and (d) U(VI) in the column effluent (C, open symbols) and well bore (W, closed symbols) throughout the experiment. The column represents an extension of the amended aquifer, and no clear difference between C and W values is evident.

(2006) who showed S^0 -dependent autotrophic denitrification is 10 times slower than heterotrophic denitrification, and the rate of denitrification with sulfide is twice as fast as with S^0 . Comparisons were also made using an equal rate for autotrophic and heterotrophic pathways.

A half-saturation constant (that is, substrate concentration, where the specific growth rate equals half its maximum rate) of $2.1 \mu\text{M NO}_3^-$ ($0.03 \text{ mg l}^{-1} \text{ NO}_3^- \text{-N}$, determined for the reaction $S^0 \rightarrow$ sulfate; Batchelor and Lawrence, 1978) was used for the autotrophic denitrification pathway. This is close to a value, $3 \mu\text{M}$, determined by Claus and Kutzner (1985), and to the values reported for aerobic sulfide oxidation (Sorokin *et al.*, 2003). A range of half-saturation constants, 11.4 – $142 \mu\text{M NO}_3^-$ (0.16 – $2 \text{ mg l}^{-1} \text{ NO}_3^- \text{-N}$), were used for the heterotrophic pathway-based values from the literature determined with either sludge or methanol as an organic carbon source (Engberg and Schroeder, 1975; Æsøy and Ødegaard, 1994; Henze *et al.* 2000; Klas *et al.*, 2006; Sarioglu *et al.*, 2009).

The catabolic reactions considered in the modeling are:



with log equilibrium constants of 3.8305, 18.2813, 13.5785 and 19.1559, respectively.

Accession numbers

Reconstructed and cloned 16S rRNA gene sequences have been deposited in GenBank under the accession numbers JX125436–JX125454 and JX120370–JX120502, respectively. Metagenomic data are accessible through DDBJ/EMBL/GenBank (SRX149542 and AMQJ00000000 (version one: AMQJ01000000), BioProject PRJNA167727), and at <http://genegrabber-berkeley.edu/2009RifleSedimentMetagenome/>.

Results and Discussion

Community composition and biogeochemical response to amendment

Previous studies have documented a period of Fe(III) reduction, followed by sulfate reduction in the Rifle aquifer during acetate amendment experiments, and concomitant reduction of U(VI) (Anderson *et al.*, 2003; Williams *et al.*, 2011). Periods marked by Fe(III) and sulfate reduction were accompanied by enrichments of *Desulfuromonadales*, and then *Peptococcaceae* followed by *Desulfobacterales* (Anderson *et al.*, 2003; Handley *et al.*, 2012). A legacy effect in the stimulated

portions of the aquifer was observed to cause earlier onset of sulfate reduction upon subsequent re-stimulation, reducing the time from >1 month to <1 week over three subsequent experiments (Callister *et al.*, 2010; Williams *et al.*, 2011; Druhan *et al.*, 2012). In this study (a third-time stimulation experiment), the onset of sulfate reduction occurred within 2 days of acetate being detected in the incubation well (Figure 1a). Sediment was recovered after at least 18 days of sulfate reduction, during which time acetate remained in excess, and U(VI)_(aq) and sulfate concentrations were halved (Figure 1b and d). Sulfide accumulation corresponded to the depletion of aqueous sulfate and Fe(II) (Figure 1c). The color of sediment throughout the column transformed from brown to black owing to the precipitation of FeS.

Clone and metagenome-derived (that is, EMIRGE-reconstructed) 16S rRNA gene data revealed a less even, lower diversity community after amendment (Figure 2a), consistent with earlier deeply-sampled 16S rRNA microarray analyses that measured $\sim 40\%$ lower OTU richness in the amended *versus* un-amended sediment (Handley *et al.*, 2012). Amendment enriched primarily for *Delta*- and *Epsilon*-*proteobacteria*, in particular a *deltaproteobacterium* closely related to the acetate-oxidizing SRB *Desulfobacter postgatei* (99% 16S rRNA gene sequence identity; Widdel and Pfennig, 1981). Other less-enriched bacteria are related to *Sulfurovum* and *Sulfurimonas*, *Geobacter* and *Desulfuromonas* and *Bacteroidetes* (Figure 2b). In general, three methods used for determining community composition and/or abundance (clone library, EMIRGE, read coverage) agreed (Supplementary Figures S2 and S3). The metagenome comprises a sample of highly abundant microorganisms; no archaeal 16S rRNA genes were detected.

Although previously un-amended sediment was used in the column, the community composition in both the column sediment, and in a split-phase sediment–quartz column incubated for the same period in well P104, differed substantially from compositions observed in first-time stimulation experiments after a similar period of amendment (Anderson *et al.*, 2003; Handley *et al.*, 2012). Less difference was observed between the column communities and those from second-time experiments, including post-amendment sediment collected from P104 in the previous year (Handley *et al.*, 2012). This tends to suggest the sediment (and quartz) were influenced by the local, twice previously stimulated, aquifer community.

Assembled genomes

After assembling Illumina reads (Table 1), we obtained genomes from seven different phylogenetic groups (Figure 3), and allocated 99% of sequence length >2 -kb long to a genomic bin (Figure 4a). Phylogenetic analyses indicate that the binned

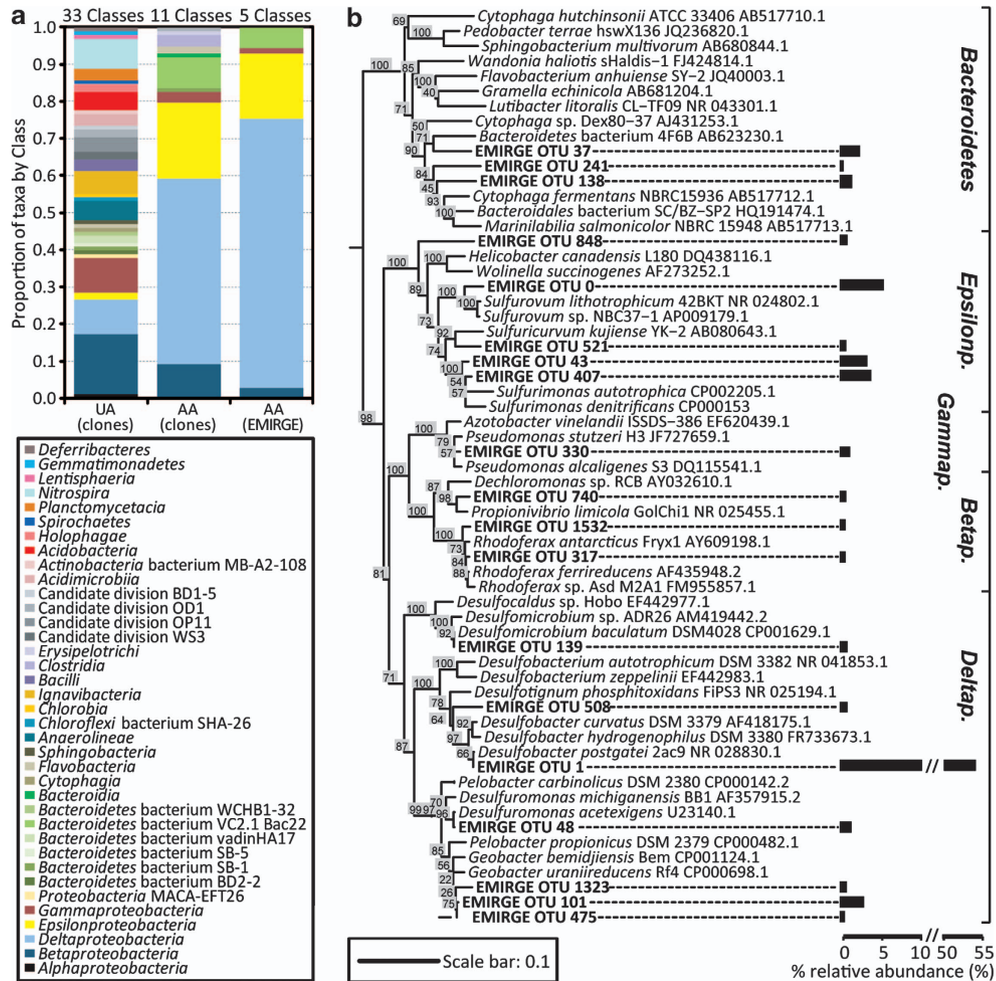


Figure 2 Community composition. (a) Relative abundances of Classes in the un-amended (UA) and acetate-amended (AA) communities determined using 16S rRNA gene clone libraries (UA and AA) or EMIRGE-reconstructed 16S rRNA genes from the metagenome (AA). (b) Bootstrap consensus tree of EMIRGE-reconstructed 16S rRNA gene sequences (AA). The tree is rooted to *Methanococcus vannielii* SB (CP000742.1). The scale bar (LHS) indicates the number of nucleotide substitutions per site. Bootstrap values are shown as percentages. A similar topology was obtained with the initial ML tree. Black bars indicate the OTU relative abundances in the metagenome. *Proteobacteria* classes are abbreviated.

genomes belong to: *Deltaproteobacteria* (r9c1, r9c7-r9c8), *Epsilonproteobacteria* (r9c2-r9c3), *Bacteroidetes* (r9c4-r9c5) and *Firmicutes* (r9c6) (Table 2). All the genomes, except for the low-coverage *Firmicutes* genome, are represented by EMIRGE-reconstructed 16S rRNA sequences. Clone library data indicate the most abundant *Firmicutes* in the amended sediment were *Clostridiales* bacteria related to *Gracilibacteraceae* or *Peptococcaceae* (data not shown). However, greater similarity to a representative *Clostridiales* genome as opposed to *Peptococcaceae* is indicated by the average amino-acid identities of orthologs (Table 2).

We estimate that the seven major genomic bins comprise ~6 near-complete and three at least half-complete genomes (Figure 4b), of which, r9c1 and r9c2 each contain a single near-complete genome, with the latter being in two PE-scaffolds (1.9 and 0.1-Mb long). R9c3 and r9c5 each contain at least two near-complete genomes. The r9c7 bin contains partial-genome fragments closely related to

Geobacter and *Desulfuromonas*. Similarity between these bacteria in terms of phylogeny, GC-content and abundance levels makes it difficult to resolve r9c7 genomes into separate bins.

Ortholog identities of > 85% have been shown to correlate with DNA-DNA hybridization values of > 70%, indicative of species level assignment (Goris *et al.*, 2007). On the basis of very high (96%) ortholog average amino-acid identity shared between r9c1 and *D. postgatei*, we suggest that they belong to same species. Excluding the small collection of r9c8 genes, which are very closely related to *Desulfomicrobium baculatum*, the other genomes have identities too low to sequenced genomes to suggest species assignment (Table 2).

Proteogenomics

A total of 1695 proteins and 112 606 peptides (93% unique peptides) from the acetate-amended samples were identified by liquid chromatography-

Table 1 Summary statistics for the Illumina Velvet and Velvet-Columbus assemblies, and the 454 Newbler assembly

Parameters	Illumina + Velvet ^a	Illumina + Columbus ^a	Newbler + 454 ^b
PE-scaffolds/contigs (# >500 bp)	11 622	10 499	17 963
N50 (kb, >500 bp)	7.7	35.2	1.8
Summed length (Mb)	48	58	27
Average length (kb)	3.8	5.6	1.5
Longest PE-scaffold/contig (kb)	114	1910	90

Abbreviation: PE, paired-end.

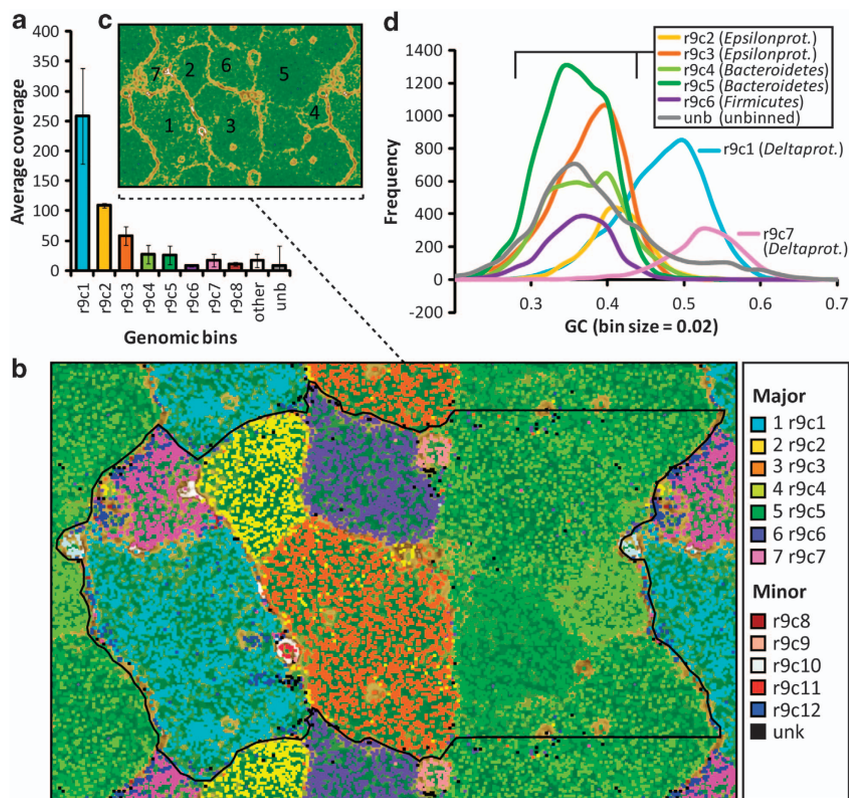
^aIllumina contigs are scaffolded using paired-end information into supercontigs (PE-scaffolds), and contain Ns.^b454 contigs were ordered (but not joined) into 2046 scaffolds using mate-pair reads.

Figure 3 Genomic bins from the Illumina assembled data distinguished by (a) average read coverage per PE-scaffold ('other' includes bins r9c8-r9c12 and unk; 'unb' comprises unassigned fragments 500–2000 bp long), (b, c) ESOM (repeating tiled view—the black line demarks one representation of each genomic bin), and (d) GC content per gene. In the ESOM, several major genomic bins occur as distinct clusters demarcated by topographic highs (beige lines, c). Two minor fragmentary genomic bins (r9c8 and r9c12) were not substantial or distinct enough from clusters 1 and 7 to form separate clusters in the ESOM, but both group with the *Deltaproteobacteria*. The three small ESOM clusters, r9c9–r9c11, are putative plasmid sequences, characterized by genes associated with transmembrane regions, DNA polymerase, phage, coiled-coil proteins, and a large number of hypothetical genes. 'Unk' are phylogenetically unassigned fragments >2 kb long. (c) ESOM without data points. Dark green denotes areas of high similarity. *Epsilonprot*, *Epsilonproteobacteria*; *Deltaprot.*, *Deltaproteobacteria*.

tandem mass spectrometry using the metagenome as a protein reference database (Supplementary Dataset). Little digestible protein was present, and few (~31) proteins were identified in the unamended sediment after searching peptides against the same database (Supplementary Information), and are not discussed further.

Examination of organism-resolved proteins enabled detection of six bacterial groups (r9c1-r9c5 and r9c7) active during the amendment experiment (Figure 5; Supplementary Information/Data set). All

groups expressed proteins associated with acetate utilization or carbon fixation *via* the TCA cycle. Among other key enzymes detected were those used in ribosomes, cell division, glycolysis/gluconeogenesis, acetate/acetyl-CoA formation, electron transport, nitrogen fixation, denitrification, sulfur respiration, phage resistance, chemotaxis and motility and response to oxidative stress. The majority of proteins detected (75% of proteins) and protein expression levels (95% of peptide spectral counts) are attributable to the dominant r9c1 genome.

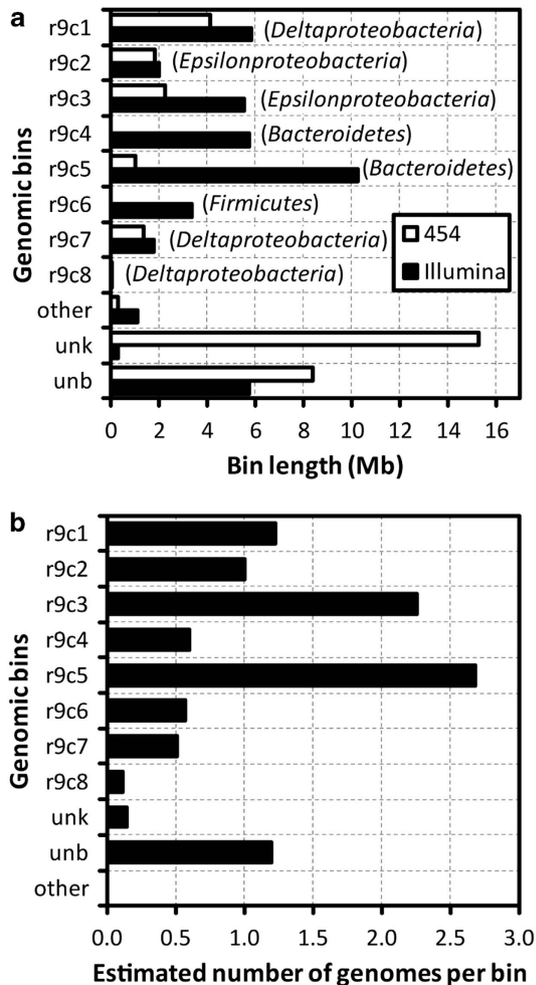


Figure 4 Plots showing (a) the length of sequence in Illumina- and 454-based genome bins; and (b) the estimated number of genomes per bin based on BLASTP searches for conserved OGs (bin lengths divided by estimated genome lengths). The sum of OGs detected in each bin was divided by the number of OGs queried (i.e. 35 *via* BLAST). ‘Other’ comprises bins r9c9–r9c12. ‘Unb’ comprises unassigned fragments 500–2000bp long that contain predicted genes (‘unk’ are unassigned fragments >2 kb). Comparisons largely agree between estimates based on OGs and those determined based on the genome size of closely related organisms ($R^2 = 0.86$; Supplementary Figure S4) even though genome sizes of related organisms can only be considered as an approximate guide.

Activity and genomic potential of enriched bacteria

Acetate-oxidizing SRB. Amending the system with acetate created a niche ecosystem for *Desulfobacter* (r9c1), which proteomic data indicate actively respired sulfate coupled to acetate oxidation (Figures 5a–c). *Desulfobacter* species (almost) dedicatedly couple anaerobic acetate oxidation to the reduction of sulfate, and were originally considered marine bacteria, before the isolation of a freshwater sediment strain (Widdel and Pfennig, 1981; Widdel, 1987). Within the Rifle aquifer, acetate-stimulated enrichments of *Desulfobacteraceae* appear to supersede enrichments of sulfate-reducing *Peptococcaceae* (Anderson *et al.*, 2003; Handley *et al.*, 2012),

although the reason for this is currently undetermined.

In this study, key proteins essential for the activation and reduction of sulfate were only detected for r9c1 (Figures 5 and 6a), including proteins potentially involved in the transfer of electrons from the membrane to APS and sulfite (QmoABC and HmeCDE; Mander *et al.*, 2002; Pires *et al.*, 2003; Mussman *et al.*, 2005). Genes were identified for *sat1* and *hmeB*, but were not represented in the proteome. In addition, almost all proteins necessary for gluconeogenesis were detected (Supplementary Information/Data set). Proteomic data (Figure 5) further indicate that r9c1 was actively dividing; chemotactic; motile by means of flagella and twitching pili; uptaking ammonium; and fixing nitrogen, possibly assisted by Rnf in electron transport (Schmehl *et al.*, 1993). Nitrogenase and ammonium transporter *amtB* have been shown to be co-expressed by acetate-stimulated blooms of *Geobacteraceae* in the Rifle aquifer during peak Fe(III) reduction (Mouser *et al.*, 2009). Expression of *amtB* transporter (and sensor) increases as ammonium concentrations approach zero (Javelle *et al.*, 2004; Mouser *et al.*, 2009) and nitrogen fixation becomes feasible (Helber *et al.*, 1988; Holmes *et al.*, 2004). While the high relative abundance of r9c1 was probably supported by nitrogen fixation, this seems unlikely to have conferred a substantial competitive advantage to r9c1 over other resident SRB (Handley *et al.*, 2012)—many of which are associated with genera containing diazotrophs (Zehr *et al.*, 2003).

We detected all but one r9c1 protein needed to oxidize acetate using a modified TCA cycle (Table 3). This cycle is used by *Desulfobacter* species instead of the acetyl-CoA pathway, which is used by other acetate-oxidizing SRB (Thauer *et al.*, 1989). The modified cycle makes use of 2-oxoglutarate:ferredoxin oxidoreductase and a reversible citrate lyase, which generate reduced ferredoxin instead of NADH and an extra ATP, respectively (Brandis-Heep *et al.*, 1983; Müller *et al.*, 1987; Müller-Zinkhan and Thauer, 1988). Neither the expected membrane-bound malate:quinone oxidoreductase (EC:1.1.5.4; Brandis-Heep *et al.*, 1983) nor the typical TCA cycle malate dehydrogenase (NADH-forming) was identified in the r9c1 genome by homology searches. While this could be an artifact of genome incompleteness, divergent homology or an unidentified analog, proteomics identified a pyruvate-forming malate dehydrogenase (NAD⁺/NADP⁺), which may potentially bypass this step, and regenerate oxaloacetate *via* pyruvate carboxylase or phosphoenolpyruvate (PEP) (cf. Hansen and Juni, 1979). Proteins for succinyl-CoA synthetase and for acetate activation by acetate kinase and phosphate acetyltransferase were identified by proteomics; however, identification of acetyl-CoA transferase suggests that r9c1 also used CoA transferase/hydrolase for both acetate activation and

Table 2 Summary data of genomic bins on the final Illumina Velvet–Columbus assembly

Bin	Longest PE-scaf-fold (kb)	PE-scaf-fold (#)	ARC ^a (s.d.)	ORFs (#)	Proteins (#)	Related organisms	AAIO ^b (s.d.)	16S ID ^c
<i>Deltaproteobacteria</i>								
r9c1	250	161	258 (80)	5421	1282	<i>Desulfobacter postgatei</i>	96 (7) F	99*
						<i>Desulfobacterium autotrophicum</i>	60 (14) D	89 [†]
r9c7	60	294	17 (11)	1390	23	<i>Geobacter bemidjiensis</i>	66 (18) F	98* ¹
						<i>Desulfuromonas michiganensis</i>	–	97* ²
r9c8	20	11	11 (4)	41	0	<i>Desulfuromonas acetoxidans</i>	59 (17) D	–
						<i>Desulfomicrobium baculatum</i>	88 (20) F	99*
<i>Epsilonproteobacteria</i>								
r9c2	1 910	2	109 (3)	1998	67	<i>Sulfurovum</i> sp. NBC37-1	71 (14) F	95*
r9c3	690	138	58 (15)	5678	160	<i>Sulfurimonas denitrificans</i>	79 (13) F	96*
						<i>Sulfurimonas autotrophica</i>	69 (15) F	94 [†]
<i>Bacteroidetes</i> ^d								
r9c4	530	498	27 (15)	3936	52	<i>Cytophaga hutchinsonii</i>	50 (12) F	83 [†]
r9c5	370	840	26 (16)	7321	78	<i>Cytophaga hutchinsonii</i>	50 (12) F	83 [†]
						<i>Wandonia haliotis</i>	–	88* ¹
						<i>Cytophaga fermentans</i>	–	90* ²
<i>Firmicutes</i>								
r9c6	60	395	8 (2)	1873	2	<i>Clostridium cellulovorans</i>	76 (25) F	–
						<i>Desulfosporosinus orientis</i>	52 (13) F	–
<i>Other</i>								
r9c9-r9c12	20–40	2–162	17 (11)	17–512	0–6	–	–	–

Abbreviations: ARC, average read coverage; AAIO, average amino acid identity of orthologs; D, draft reference genome; F, fully-curated reference genome; 16S ID, percent identity between 16S rRNA genes; ORFs, predicted genes; #, number.

^aARC was calculated from k-mer coverage.

^bUsing less strict criteria for determining orthologs (i.e., without a bit-score cutoff of ≥ 60) made $\leq 2\%$ difference to calculated AAIs for most of the genome bins (r9c1–r9c5), but decreased AAIs by 3–6% for r9c6 and r9c8, and by 14% for r9c7.

^cPercent identity match between EMIRGE-reconstructed and GenBank 16S rRNA sequences.

^dThe determined AAI of orthologs between r9c4 and r9c5 is 53%.

* denotes the nearest isolate sequence—*¹ and *² indicate where two reconstructed 16S rRNA gene sequences correlate to a single bin (r9c7), or cannot be assigned to one of the two bins (r9c4 and r9c5). [†] denotes matches to other organisms for comparison.

succinate formation (Figure 6b), as has been demonstrated for *D. postgatei*, *Desulfuromonas acetoxidans* and *Geobacter* species (Brandis-Heep *et al.*, 1983; Gebhardt *et al.*, 1985; Wilkins *et al.*, 2009).

In contrast to r9c1, *Desulfomicrobium* (r9c8) was less enriched. Members of this sulfate-reducing genus are incomplete oxidizers and are only known to use acetate mixotrophically with H₂ (Dias *et al.*, 2008). Assuming that r9c8 was also reducing sulfate, or other inorganic compounds respired by *Desulfomicrobium* species (for example, thiosulfate and nitrate), then it would likely have been substrate limited, and dependent on the production of H₂ or organic acids by other community members. Most *Desulfomicrobium* species are also able to ferment select organic acids (Dias *et al.*, 2008).

Other acetate oxidizers. Few proteins were identified from *Desulfuromonadales* r9c7, owing partly to incomplete genomes (Table 2, Figure 4b). Proteins and genes identified may be attributed to either the *Geobacter*- or *Desulfuromonas*-like bacteria

comprising the r9c7 bin. Some genes, but no proteins, were detected for nitrogen fixation, ATP synthase, cell division, gluconeogenesis and motility by pili or flagella, and only one protein was detected for chemotaxis. Several key TCA cycle proteins were detected (Figures 5a and d; Table 3), demonstrating that *Desulfuromonadales* were actively consuming acetate, while sulfate reduction prevailed as the major terminal electron-accepting process. These bacteria do not reduce sulfate, but are well-known for their ability to reduce metals (including iron and uranium) and other inorganic elements or compounds (Lovley *et al.*, 2004).

Studies have demonstrated a correlation between the addition of simple organic substrates, such as glucose and organic acids like acetate, to terrestrial aquifers and the enrichment of Fe(III)-reducing *Geobacter* species (Snoeyenbos-West *et al.*, 2000; Holmes *et al.*, 2002, 2007). Repeat experiments amending the Rifle aquifer with acetate have consistently produced blooms of *Geobacter* associated with Fe(III) and U(VI) reduction (for example,

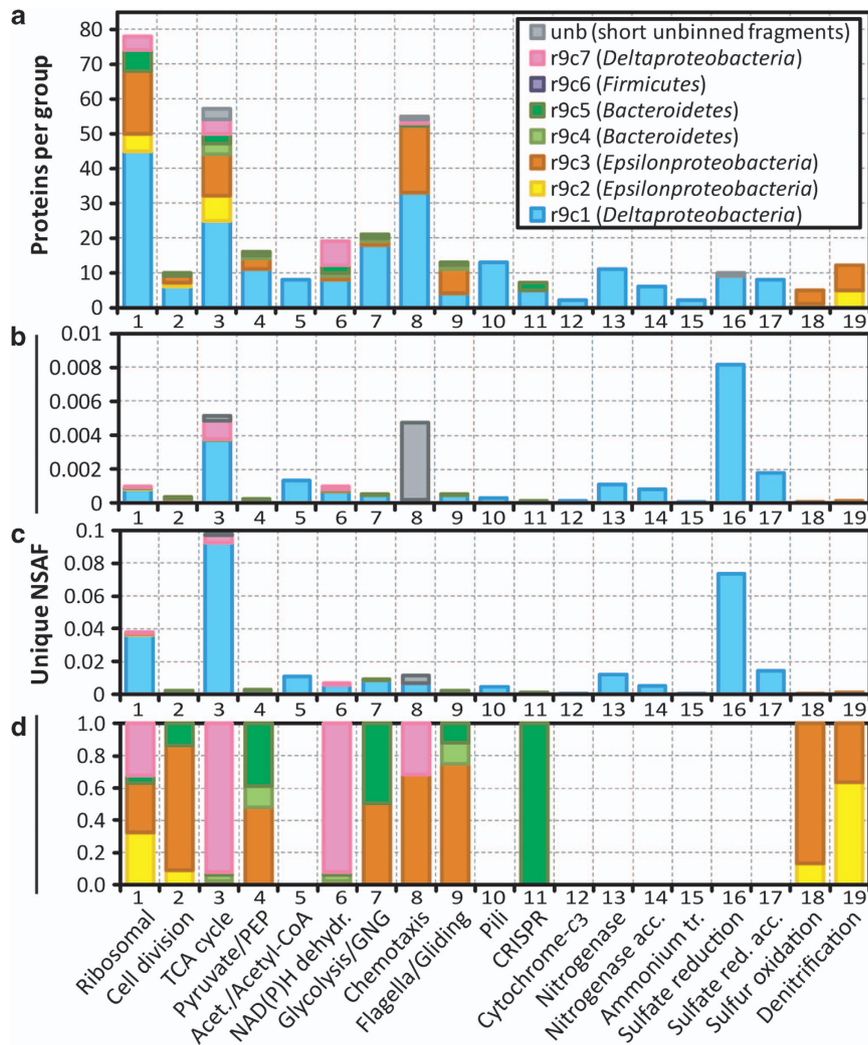


Figure 5 Proteins expressed in key functional groups (Supplementary Dataset) are shown as (a) number of proteins identified per group, (b) averaged or (c) summed unique NSAFs, and (d) relative proportions of summed unique NSAFs with r9c1 and unb removed. Little difference was evident between non-unique (not shown) and unique NSAF charts, except the overall contribution of TCA cycle peptides was 0.5% less in the latter. No proteins within these groups were matched to the r9c6 *Firmicutes* genome. Abbreviations and definitions: (3) oxidative/reductive TCA cycle proteins (4) conversion between pyruvate and phosphoenolpyruvate (PEP); (5) conversion between acetate and acetyl-CoA; (6) NADH/NADPH dehydrogenase; (7) glycolysis/gluconeogenesis; (11) Cas proteins associated with Clustered Regularly Interspaced Short Palindromic Repeats, CRISPR; (12) cytochrome-c₃ and cytochrome-c₃ hydrogenase; (13) nitrogenase (NifABDEHKU); (14) putative nitrogenase accessory proteins RnfCDEG; (15) ammonium transport AmtB; (16) sulfate reduction proteins ATP-sulfurylase (Sat2), adenylyl sulfate (APS) reductase (AprAB), dissimilatory sulfite reductase (DsrABCFEH); (17) putative accessory proteins heterodisulfide reductase-like menaquinol-oxidizing enzyme (HmeCDE) and quinone-interacting membrane-bound oxidoreductase (QmoABC); (18) sulfur oxidation proteins SoxCY and SQR; (19) NO₃ (NapAB), NO₂ (NirS), NO (NorC), N₂O (NosZ) reductases (further details in Supplementary Text and Dataset).

Anderson *et al.*, 2003; Holmes *et al.*, 2007; Wilkins *et al.*, 2009). These enriched *Geobacter* and co-enriched *Desulfuromonas* species persist as significant community members, although at lower relative abundances, after sulfate reduction becomes prevalent (Handley *et al.*, 2012), and *Geobacter* are key candidates for the ongoing reductive immobilization of U(VI) observed during sulfate reduction in this study (Figure 1d). Despite the capacity for abiotic reduction of sulfide, microbial Fe(III) reduction may proceed concurrent with sulfate reduction (Sørensen, 1982; Tugel *et al.*, 1986; Canfield, 1989).

It is therefore plausible that either genera attributed to the *Desulfuromonadales* r9c7 may have also coupled acetate-oxidation with Fe(III) reduction. Alternatively, r9c7 bacteria may have gained energy from S⁰ reduction (Pfennig and Biebl, 1976; Lovley *et al.*, 2004), which has been shown to accumulate during acetate amendment of the Rifle aquifer. Williams *et al.* (2011) identified 4–5 mmol kg⁻¹ S⁰ (below 3-m depth) in the core collected to create well P104 (used in this study) after 110 days of acetate amendment (second year amendment). Comparably, little S⁰ was identified in

Table 3 TCA cycle and related components identified for genomic bins r9c1-5 and r9c7

#	Enzyme	EC	1	2	3	4	5	7
1	Citrate synthase	2.3.3.1	—	—	G	—	P	P
1	ATP citrate lyase	2.3.3.8	P	P	P	—	—	—
1	Citrate lyase	4.1.3.6	P	G	G	G	G	—
2	Aconitate hydratase	4.2.1.3	P	G	P	—	G	G
3	Isocitrate dehydrogenase	1.1.1.41/42	P	P	P	P	G	—
4a	2-oxoglutarate dehydrogenase E1	1.2.4.2	—	—	—	G	—	—
4b	2-oxoglutarate dehydrogenase E2	2.3.1.61	—	—	—	G	G	—
4c	Dihydrolipoamide dehydrogenase	1.8.1.4	P	—	—	G	G	—
4	2-oxoglutarate synthase	1.2.7.3	P	P	P	G	G	—
5	Succinyl-CoA synthetase	6.2.1.4/5	P	P	P	G	G	—
6	Succinate/fumarate reductase	1.3.5.1 ^a	P	P	—	P	P	P
7	Fumarate hydratase	4.2.1.2	P	G	G	—	G	—
8	Malate dehydrogenase (NAD-dependent)	1.1.1.37	—	P	G	P	G	P
8	Malate dehydrogenase (quinone)	1.1.5.4	—	—	G	G	—	—
<i>Related: pyruvate/PEP</i>								
	PEP synthetase	2.7.9.2	P	G	G	G	G	—
	PEP carboxylase (GTP)	4.1.1.32	P	—	—	—	—	—
	PEP carboxylase (ATP)	4.1.1.49	—	—	—	—	—	—
	PEP carboxylase	4.1.1.31	G	—	—	—	P	—
	Pyruvate/oxaloacetate carboxylase	6.4.1.1 ^b	P	P	P	G	G	—
	Pyruvate ferredoxin oxidoreductase	1.2.7.1	P	G	P	P	—	—
a	Pyruvate dehydrogenase E1	1.2.4.1	P	G	—	G	G	—
b	Pyruvate dehydrogenase E2	2.3.1.12	P	—	—	—	—	—
c	(4c above)							
<i>Related: acetate ↔ acetyl-CoA</i>								
	Acetyl-CoA hydrolase/transferase	3.1.2.1	P	—	—	—	G	—
	Acetyl-CoA synthetase	6.2.1.1	P	G	G	—	—	—
a	Acetate kinase (acetyl-P ↔ acetate)	2.7.2.1	P	G	G	G	G	—
a	Acylphosphatase (acetyl-P → acetate)	3.6.1.7	—	G	—	G	G	—
b	Phosphate acetyltransferase	2.3.1.8	P	G	G	G	G	—

Abbreviations: #, order of TCA cycle reactions in the oxidative direction; EC, enzyme commission number; G, genes only identified; P, proteins also detected.

Reactions for enzymes not previously defined here: pyruvate carboxylase (pyruvate → oxaloacetate); pyruvate ferredoxin oxidoreductase (i.e., pyruvate synthase; pyruvate ↔ acetyl-CoA); pyruvate dehydrogenase and dihydrolipoamide dehydrogenase (pyruvate ↔ acetyl-CoA).

^aSuccinate/fumarate reductase equates to EC:1.3.5.1/EC:1.3.99.1.

^bpyruvate carboxylase equates to EC:6.4.1.1/EC:4.1.1.3.

less-stimulated sediment cores collected further from the acetate source.

Bacteroidetes, on the other hand, are well-known for their ability to degrade carbohydrates and other complex organic compounds using respiratory or fermentative metabolisms (Holmes *et al.*, 2007; Lee *et al.*, 2010; Thomas *et al.*, 2011). A number of genes identified here are associated with mannose metabolism (mannose-1-phosphate guanyltansferase, mannose-6-phosphate isomerase, GDP-mannose 4,6-dehydratase, phosphomannomutase), xylan degradation (candidate β -xylosidase, endo-1,4- β -xylanase) and cellulose degradation (family 3 candidate β -glycosidase), including identification of *Bacteroidetes*-like unbinned partial fragments (0.5–2 kb long) resembling candidate β -glycosidase and cellobiose phosphorylase. However, few expressed proteins indicative of these metabolisms were identified. Only two proteins (with peptide counts just above detection) were identified that are suggestive of cellobiose degradation and mannose metabolism (candidate β -glycosidase hydrolase, phosphomannomutase), and only two glycolysis proteins were detected.

Instead, detection of some key TCA cycle enzymes, including citrate synthase (Figures 5a and d; Table 3), suggests that the *Bacteroidetes* may have been using the cycle heterotrophically to oxidize acetate (cf. Xie *et al.*, 2007; Zhang *et al.*, 2009), although use of the reductive cycle cannot be excluded. Genomic data indicate that *Bacteroidetes* r9c5 has a complete oxidative TCA cycle (near-complete in r9c4), and both r9c4 and r9c5 have genes that may support a reductive cycle (that is, citrate lyase, 2-oxoglutarate synthase, pyruvate synthase, PEP synthetase and carboxylase). The electron acceptor for this reaction is not evident; however, on the basis of genomic evidence, one possibility is that *Bacteroidetes* coupled acetate oxidation to the reduction of nitrogen species, as genes were identified for nitric-oxide reductase (*norABD*) and nitrous-oxide reductase (*nosLZ*).

Genes were also identified for a formate-dependent, ammonium-forming nitrite/(polysulfide) reductase (*nrfADH*); selenate reductase *ygfK* (r9c4 only); sulfur metabolism, namely SQR (r9c4 only), a *Chlorobium luteolum*-like flavocytochrome-*c* sulfide dehydrogenase and a *phsC*-like

cytochrome-*b*₅₆₁ thiosulfate reductase; and arsenic detoxification (similar to r9c2-r9c3; Figure 6a). Although the experimental condition under which *Bacteroidetes* r9c4 and r9c5 were growing was anoxic, genes for cytochrome-*c* and cytochrome-*bd* oxidase and cytochrome-*cbb*₃ oxidase (*ccoPQNOS*) indicate a capacity for these bacteria to also grow (micro)aerobically (Preisig *et al.*, 1993; Trumpower and Gennis, 1994; Visser *et al.*, 1997; Kulajta *et al.*, 2006).

Sulfur oxidizers: acetate-induced syntrophy. Previously, researchers have demonstrated syntrophic growth between oxygen-dependent, sulfide-oxidizing *Thiobacillus thioparus* and sulfate-/S⁰-reducing, *Desulfovibrio desulfuricans* in co-culture, such that a positive-feedback cycle was established between the reductive and oxidative processes of these two bacteria (van den Ende *et al.*, 1997). In another study, geochemical and functional gene-based evidence for a cryptic microbial sulfur cycle, suggests that this process may have important biogeochemical consequences for sulfate recycling in oceanic oxygen minimum zones (Canfield *et al.*, 2010). Proteomic results here indicate that a similar process was operating in the acetate-amended Rifle aquifer sediment, and suggest that syntrophic growth of autotrophic or possibly mixotrophic *Sulfurovum*- and *Sulfurimonas*-like bacteria (r9c2 and r9c3) were supported by sulfide and CO₂ generated by heterotrophic respiration.

Species within these two epsilonproteobacterial genera are autotrophic, and oxidize sulfide, sulfur and thiosulfate either with oxygen and/or nitrate serving as electron acceptors (Hoor, 1975; Inagaki *et al.*, 2003, 2004). *Sulfurimonas denitrificans* oxidizes sulfide by reducing nitrate completely to N₂ (Hoor, 1975). According to genomic studies of *Sulfurovum* sp. NBC37-1 and *S. denitrificans* DSM1251, sulfur oxidation in these organisms may proceed *via* the sulfur oxidation pathway (SoxCDYZXAB), forming sulfate, or SQR, forming S⁰ from sulfide (Nakagawa *et al.*, 2007; Sievert *et al.*, 2008).

Similarly, r9c2 and r9c3 possess respiratory genes for a complete denitrification pathway (*napABDFGHL*, *nirSF*, *nirNJ* for r9c3 only, *norBC*, *nosZ*), sulfur oxidation (SQR, *soxCDYZ*) excluding *soxXAB* (Supplementary Figure S5), and also the reductive TCA cycle for CO₂ fixation (Table 3; Figure 6b). Proteomic data infer that, during amendment these *Epsilonproteobacteria* performed nitrate-dependent sulfide/sulfur oxidation coupled to CO₂ fixation *via* the reductive TCA cycle (Figures 5a and d). However, simultaneous operation of an oxidative TCA cycle to support mixotrophic growth with acetate cannot be excluded (cf. Tang and Blankenship, 2010). Proteomics further show r9c2 and r9c3 were dividing, and r9c3 was chemotactic and motile. Genes for cytochrome-*cbb*₃ oxidase (*ccoPQONS*), cytochrome-*b*₅₆₁ (r9c3; Murakami

et al., 1986) and cytochrome-*bd* (r9c2) suggest that these bacteria could also oxidize sulfur (micro)aerobically.

The missing *soxXAB* genes (not detected by homology searches) are normally required for thiosulfate (and presumably sulfide and sulfur) attachment to SoxYZ, activation to sulfane, and release after oxidation by SoxCD (Friedrich *et al.*, 2005; Sauv   *et al.*, 2007; Zander *et al.*, 2011). Hence, the exact function of Sox in sulfur oxidation by r9c2 and r9c3 cannot be deduced from our data. Green sulfur bacteria lacking *soxCD* (but with functional *soxXAB* and *soxYZ* genes) are able to use *dsr* with or without *apr* genes in reverse to oxidize sulfide or sulfite, respectively (Sakurai *et al.*, 2010; Gregersen *et al.*, 2011). We identified no genes indicative of a reverse sulfate reduction pathway in r9c2 and r9c3. Nevertheless, *in vitro* enzyme assays by Rother *et al.* (2001) suggest sulfide or S⁰ oxidation may proceed, although at a slower rate (13–19 or 3–17 times less, respectively), without either SoxXA or SoxB. The same is true for SoxXAB without SoxCD or SoxYZ.

Synteny is shared between *soxXAB* and *soxCDYZ* in the model organism, *Paracoccus pantotrophus*, and several other *Alphaproteobacteria* (Friedrich *et al.*, 2005). However, they form two non-synteny gene clusters in bacteria closely related to r9c2 and r9c3, *Sulfurovum* sp. NBC37-1 and *S. denitrificans* DSM1251 (Nakagawa *et al.*, 2007; Sievert *et al.*, 2008; Supplementary Figure S5). In as much as conservation of gene order tends to suggest conservation of gene function, loss of synteny tends to suggest a loss of co-dependence between the gene clusters, possibly occurring with increased evolutionary distance (Yelton *et al.*, 2011). This may explain the apparent loss of *soxXAB* genes in r9c2 and r9c3.

While it is possible that r9c2 and r9c3 completely re-oxidized sulfide to sulfate, an autotrophic denitrifying community from anaerobic sludge has been shown to only partially oxidize sulfide (probably to S⁰) when placed under nitrate-limiting conditions, but completely oxidize sulfide to sulfate with unlimited nitrate (Cardoso *et al.*, 2006). Considering the micromolar concentrations of nitrate available in the aquifer (Williams *et al.*, 2011), this could also reasonably explain the enrichment of bacteria in this study (r9c2 and r9c3) that are potentially only able to oxidize sulfide to S⁰. Any formation of S⁰ by these bacteria would likely be re-cycled by putative sulfur-reducing bacteria, r9c7.

Autotrophic versus heterotrophic denitrification

Simulations of biogeochemical processes occurring during acetate amendment captured the general features of the acetate and sulfate breakthrough curves at well P104, and estimate nitrate conversion to N₂ (Supplementary Figures S6a and b). Although simulations for autotrophic (sulfide-dependent) and heterotrophic (acetate-dependent) denitrification

use estimates for nitrate and specific growth rates, and exclude biomass, analyses indicate that the autotrophic denitrification pathway, involving sulfide oxidation to S^0 or sulfate, is thermodynamically feasible. Despite its greater standard Gibbs energy of reaction, the autotrophic sulfide to sulfate pathway is less favored than the sulfide to S^0 pathway (Supplementary Figure S6c) because of the high sulfate concentration in groundwater (Figure 1b).

Although heterotrophic denitrification is thermodynamically favored over autotrophic denitrification (ΔG of ca. -220 versus -178 kJ per electron per mole), the effect is insignificant if the thermodynamic factor (F_T) formulation in Equation 3 (Supplementary Information; Supplementary Figure S6c) is assumed to be correct, as the system is effectively far from equilibrium with respect to both pathways, and F_T is very close to 1. This implies that behavior is controlled by the Monod kinetic terms in Equation 2 (Supplementary Information). Principally owing to the low half-saturation constant used for autotrophic denitrification, simulations for the sulfide–sulfur reaction tend to indicate that the autotrophic pathway dominates over the heterotrophic pathway given a concentration of $5\ \mu\text{M}$ (at all considered heterotrophic versus autotrophic rates), and vice versa for the higher $72\text{-}\mu\text{M}$ concentration when assuming a slower autotrophic rate (Supplementary Figure S6d–i). While a potential impact of sulfide toxicity on community composition and function cannot be excluded, modeling results may explain why, with acetate in excess, autotrophic sulfide-dependent denitrification by *Epsilonproteobacteria* r9c2 and r9c3 out-competed heterotrophic denitrification in this experiment. Another factor may be possible mixotrophic growth by *Epsilonproteobacteria* r9c2 or r9c3, which would likely increase cell growth and denitrification rates (Cardoso *et al.*, 2006).

Conclusions

We reconstructed the genomes of members of a subsurface sediment community enriched during acetate amendment. Proteomics identified organism-specific function and syntrophic interactions among community members (Figures 6a and c). While the dominant process identified was acetate-fueled sulfate reduction, excess acetate was also respired by enriched *Desulfuromonadales*, probably supporting concurrent Fe(III)-, U(VI)- and/or S^0 -reduction. Gene-based evidence and TCA cycle proteins detected for *Bacteroidetes* suggest that they contributed to acetate degradation, and have the capacity for reducing nitrogen species. Co-enrichment of *Epsilonproteobacteria* and expression of proteins associated with sulfide oxidation and carbon fixation imply that products of heterotrophic acetate oxidation—sulfide and CO_2 —were used as a carbon and energy source by autotrophic or

mixotrophic *Epsilonproteobacteria*. Sulfide-dependent denitrification may have been favored over the heterotrophic pathway owing to nitrate-limiting conditions within the Rifle aquifer, such that kinetic factors govern outcomes. In turn, reaction products of epsilonproteobacterial metabolism, such as N_2 , were probably fixed and sulfate or S^0 respired by *Desulfobacter* or *Desulfuromonadales* bacteria. These results suggest acetate-amendment promoted complex organismal and metabolic processes and interactions involved in carbon, sulfur, metal and nitrogen cycling.

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