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Hexavalent Chromium Reduction under Fermentative Conditions with Lactate Stimulated Native Microbial Communities

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Abstract

Microbial reduction of toxic hexavalent chromium (Cr(VI)) in-situ is a plausible bioremediation strategy in electron-acceptor limited environments. However, higher [Cr(VI)] may impose stress on syntrophic communities and impact community structure and function. The study objectives were to understand the impacts of Cr(VI) concentrations on community structure and on the Cr(VI)-reduction potential of groundwater communities at Hanford, WA. Steady state continuous flow bioreactors were used to grow native communities enriched with lactate (30 mM) and continuously amended with Cr(VI) at 0.0 (No-Cr), 0.1 (Low-Cr) and 3.0 (High-Cr) mg/L. Microbial growth, metabolites, Cr(VI), 16S rRNA gene sequences and GeoChip based functional gene composition were monitored for 15 weeks. Temporal trends and differences in growth, metabolite profiles, and community composition were observed, largely between Low-Cr and High-Cr bioreactors. In both High-Cr and Low-Cr bioreactors, Cr(VI) levels were below detection from week 1 until week 15. With lactate enrichment, native bacterial diversity substantially decreased as Pelosinus spp., and Sporotalea spp., became the dominant groups, but did not significantly differ between Cr concentrations. The Archaea diversity also substantially decreased after lactate enrichment from Methanoseta (35%), Methanosarcina (17%) and others, to mostly Methanosarcina spp. (95%). Methane production was lower in High-Cr reactors suggesting some inhibition of methanogens. Several key functional genes were distinct in Low-Cr bioreactors compared to High-Cr. Among the Cr resistant microbes, Burkholderia vietnamiensis, Comamonas testosterone andRalstonia pickettii proliferated in Cr amended bioreactors. In-situ fermentative conditions facilitated Cr(VI) reduction, and as a result 3.0 mg/L Cr(VI) did not impact the overall bacterial community structure.

Introduction

Subsurface heavy metal contamination from the nuclear weapons industry is a continuing problem at the Department of Energy site at Hanford, WA. Hexavalent chromium (Cr(VI)) is toxic and highly soluble, and, as a result, can be readily transported through the groundwater [1]. When Cr(VI) is reduced to Cr(III), the solubility and mobility decrease [2], except when Cr(III)-organic complexes can also become soluble and toxic [3]. Microbial Cr(VI)-reduction is one plausible remediation strategy for contaminated sites, with a wide diversity of microorganisms identified to be capable of reducing Cr(VI) as well as other metals [2,4,5]. Sustainable long-term microbial Cr(VI)-reduction can be challenging as it is dependent upon several biotic and abiotic processes including physiological, hydrological and geochemical parameters that subsequently control the stability of Cr(III) [6]. Additionally, in-situ microbial metal-reduction over time can become inefficient by limited supply of electron donors and acceptors [7,8,9].
As an example, polylactate hydrogen release compound (pHRC) was developed and evaluated for overcoming some issues of long term Cr(VI)-reduction for in-situ remediation of Cr(VI) contaminated groundwater at the Hanford site [1,10]. Microcosm experiments stimulated the indigenous microbial community but also shifted the community composition that was able to reduce Cr(VI) [10]. The in-situ injection of pHRC into groundwater also stimulated the native microbial community and led to a depletion of higher redox terminal electron acceptors, increasing Fe(II) and significantly decreasing Cr(VI) [1]. Hence, it appeared that in the presence of excess electron donors, the microbial populations may have become more specialized.

Depending upon the type of electron donors and availability of terminal electron acceptors, the stimulated microbial community could be dominated by syntrophs and fermenters, such as Pelosinus spp. with lactate enrichment as evidenced in our previous study [11]. Pelosinus spp. outcompeted sulfate- and Fe(III)- reducers in a terminal electron limiting environment and some of the species have been shown to reduce variety of metals including Cr(VI) [11,12], that could be stimulated for metal reduction. Fermentative and methanogenic conditions involve complex syntrophic interactions for stable community structure and function, but not much is known about the impacts of Cr(VI) concentrations on long term stability of fermentative/syntrophic/methanogenic communities and reduction of Cr(VI). It is well established though, that Cr(VI) can be toxic to variety of microorganisms [13,14]. We anticipated that with exposure to varying levels of Cr(VI), indigenous microorganisms during longer-term lactate enrichment will shift and select for Cr-resistant syntrophic communities. The goals of this study were to determine the lactate enriched native microbial community response to different levels of Cr(VI) concentrations as well as the subsequent Cr(VI)-reduction rate and extent. Continuous flow bioreactors were implemented to enrich native communities from a contaminated site at Hanford, WA with three Cr(VI) concentrations representing the uncontaminated area (0.0 mg/L), the edge of the encroaching Cr plume (0.1 mg/L) and the center of the Cr plume (3.0 mg/L) at the site.

**Materials and Methods**

**Native community cultivation and monitoring**

Groundwater samples were collected from well 699-96-43 at a depth of 42.5 feet, which is <1 foot from the Hanford formation bedrock, at 100H at the Department of Energy’s Hanford Site (Washington, USA) [1]. A description of the site contamination and geochemical history has been previously described [1,15]. Briefly, Cr(VI) contamination likely emanates from sodium dichromate used for corrosion control at the Hanford plutonium reactors. Cr(VI) groundwater concentrations decrease from the source (area 100D, ~3.0 mg/L Cr(VI)) to near the shore of the Columbia river (area 100H) at ~0.1 mg/L Cr(VI). Samples (600 ml) were sealed under N2, placed on blue ice packs and shipped to Oak Ridge National Laboratory. Upon arrival, 50 ml was removed as a reference, and immediately frozen at ~80°C. The remaining groundwater (550 ml) was inoculated into six custom-built, anaerobic glass fermentation vessels as described previously [11] and in the Supplemental Materials and Methods.

Each vessel received 90 ml of ground water, with working volumes of ~800 ml with the growth media. Modified CCM media [16] with 30 mM sodium lactate but without exogenous electron acceptors was used for enrichment of the culture (more details are provided in Supplemental Materials and Methods). The duplicate treatments were established where bioreactors 1 and 2 received 30 mM lactate and 0.1 mg/L Cr(VI) (Low-Cr), bioreactors 3 and 4 received 30 mM lactate enrichment only (No-Cr) and bioreactors 5 and 6 received 30 mM lactate and 3.0 mg/L Cr(VI) (High-Cr). The Cr(VI) was added in the form of Na2Cr2O7. The duplicate bioreactors were each supplied with medium from one of three 19 L carboys. Outflow samples from each bioreactor were taken bi-weekly for monitoring growth (OD600), pH, Cr(VI), lactate, acetate and butyrate. Growth within the bioreactors was compared to an equal volume taken from the sterile medium carboys. Gas samples were taken aseptically with needles and syringes through vessel top ports sealed with butyl rubber stoppers to measure CO2, H2 and CH4. A separate media outflow sample was taken weekly for Cr(VI) and Fe(III) reduction potential assays. Cr(VI) concentrations were also measured weekly in the samples taken from the “feed carboys” to detect any abiotic reduction of Cr(VI).

**Metabolites, Cr(VI) and metal reduction monitoring in bioreactor samples**

Lactate, acetate and butyrate from the media samples were measured using a Waters Breeze 2 HPLC system (Waters Corp., USA). Headspace samples were injected to a Agilent 6850 GC (Agilent Technologies, USA) equipped with a thermal conductivity detector (TCD) for CO2 and H2 quantification. Samples to detect CH4 concentrations were injected into an Agilent 6890 gas chromatograph equipped with a flame ionization detector (FID).

Samples from bioreactors were centrifuged (13,000 xg, 8 min, 4°C), cell pellets were washed 3 times with 30 mM lactate/30 mM NaHCO3 buffer (pH 6.8) and finally resuspended to 8 ml with the 30 mM lactate/30 mM NaHCO3 buffer [17] for Cr(VI) and Fe(III) -reduction assays. The diphenylcarbazide [18] and the ferrozine methods [19], respectively, were used for measuring Cr(VI) and Fe(III) concentrations from the media and resuspended cell pellet samples. Further details on metabolites, and metal-reduction assays are given in the Supplementary Materials and Methods.

**DNA extraction and pyrosequencing of the SSU rRNA subunit of Bacteria and Archaea**

For pyrosequencing of the SSU rRNA subunit of Bacteria and Archaea, 13 ml media samples were collected every week from bioreactor outflows, centrifuged and pellets were stored at ~80°C until analysis. Total community genomic DNA (cgDNA) was extracted using the PowerSoil™ DNA Isolation Kit (Mo Bio Labs, Inc., Carlsbad, CA). Separate sets of primers targeting the V1–3 hyper variable regions of the SSU rRNA subunit of Bacteria and Archaea were used for PCR amplification of 16S rRNA gene. Primers used for bacteria were 27YMF (AGAGTTTGTATCMTGGCTCAG) and 534R (TYACCGCGGCTGCTGG) to get an approximate amplicon length of 431–550 bp, and A2FA (TCGSGTGATTCCYGCGS9) and 571R (GCTACGRG- YVSCCTTARRC) for Archaea for an approximate amplicon length of 479–1221 [20]. These primers were designed with FLX titanium adapters and 8–10 nt barcodes for sample multiplexing. The PCR reactions were conducted for 10 ng template in 50 µl PCR mix [21] with high fidelity AccuPrime™ Taq DNA polymerase (Invitrogen, Carlsbad, CA). The PCR amplicons were purified using the Agencourt AMPure solid-phase paramagnetic bead technology (Agencourt Bioscience Corporation, Beverly, MA). The PCR amplicon purity, concentration and size were estimated using DNA 1000 reagents and an Agilent 2100 Bioanalyzer (Agilent Technologies, Inc., Waldbronn, Germany). The reactions were paired according to DNA quantity and quality prior to performing emulsion reactions for sequencing on a 454
Life Sciences Genome Sequencer FLX (Roche Diagnostics, Indianapolis, IN) using the unidirection amplicon library sequencing protocol with emPCR Kit II (Roche) and FLX titanium chemistry.

16S rRNA amplicon sequence analysis
The 16S rRNA amplicon sequence data was analyzed in MOTHUR version 1.26 [22], QIIME (Quantitative Insights Into Microbial Ecology), version 1.5.0 [23] and AmpliconNoise V1.25 [24]. All samples were run through the AmpliconNoise pipeline to remove sequencing and PCR errors and chimeras using the built-in Perseus algorithm. Sequences were aligned in MOTHUR against the RDP (Ribosomal Database Project) databases for Bacteria and Archaea and trimmed to preserve an approximate average length of 400nt. Pairwise distances were calculated in MOTHUR, which were then clustered based on average linkage clustering. The operational taxonomic units (OTUs) were defined at 97% similarity cutoff for all analyses. BIOM format OTU table was implemented in QIIME pipeline for diversity analysis, classification and assigning taxonomy to the OTUs. Taxonomy reference database from RDP were used for assigning taxonomy to Bacteria and Archaea OTUs. Pairwise Bray-Curtis distances between samples for the OTU table without singletons were used to create non-metric multidimensional scaling plots using the PAST software package [25].

GeoChip microarray hybridization
Bioreactor DNA samples from selected time points (0.5 w (weeks), 3 w, 6 w, 12 w and 15 w) were submitted to the Institute of Environmental Genomics at Oklahoma University, Norman, OK for GeoChip microarray hybridization. GeoChip 4.0 was used for hybridization. This version contains approximately 132,000 probes covering 410 gene categories [26]. DNA labeling, hybridization, initial data normalization and processing parameters are as previously described [26]. For comparative analysis among the samples, we further normalized the data by log transforming the gene probe signal intensity. NMDS plots for Bray-Curtis dissimilarity distances were calculated in PAST software under default settings. Hierarchical clustering analysis and heat maps were created using heatmap.2 in Gplots package within R software [27].

Sequence accession numbers
Sequences from this study were deposited in the GenBank Short Read Archive database under accession number SRA072678.

Results
Microbial community growth with lactate enrichment and Cr(VI) levels
After 1 week, optical densities (OD_{600}) of the communities in all bioreactors were similar (0.39–0.52) and by 7 weeks, all bioreactors had reached densities of 0.56 or greater (Figure 1) and were maintained throughout the experiment. Notable differences were observed in the Low-Cr(VI) (0.1 mg/ml Cr(VI)) bioreactors as they showed greater fluctuation between the replicates early on but maintained a more comparable density during 9–14 weeks. It was notable that these two bioreactors also reproducibly diverged from the others at 8 weeks (0.55–0.65 vs. 0.79–1.1 OD_{600}) and it was not until 14.5 weeks that they were statistically (mean standard deviations) indiscernible from the lactate only (0.0 mg/L Cr(VI)) and High-Cr(VI) (3.0 mg/L Cr(VI)) bioreactors (0.77–0.99 OD_{600}). Reasons for this are unclear when comparing this disparity in growth from the organic acids (Figure 2) and headspace gas concentrations (Figure 3), as detailed below. Although not statistically significant, the Low-Cr bioreactors did display higher CH4 concentrations, however this trend began at 5 weeks (Figure 3B). The pH of all the bioreactors and the media carboys was 7.0±0.2 over the 105 day experiment (data not shown).

Figure 1. Averaged changes in cell densities (OD_{600}) of the microbial consortia in duplicate bioreactors inoculated with Hanford well H-100 groundwater amended with (X) 0 mg/L Cr(VI), (○) 0.1 mg/L Cr(VI) and (▲) 3.0 mg/L Cr(VI). W = weeks. Error bars indicate 1 standard deviation between replicates.
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Lactate concentrations in the spent media from all bioreactors dropped significantly from ~35 mM to ~5 mM after 1.5 weeks and became limited after 4 weeks although fluctuations were observed in individual bioreactors throughout the first 6 weeks (Figure 2A). Acetate concentrations increased to 20–30 mM in all bioreactors by week 2, and to 41–46 mM by week 4 (Figure 2B). On week 7, acetate concentrations decreased in all bioreactors to ~30 mM, but recovered and increased to 40–50 mM. Acetate concentrations in the Low-Cr reactors trended to be lower, although it was not statistically significant. Other metabolites such as butyrate and formate were analyzed, but their presence was intermittent and <2 mM (data not shown).

Headspace CO₂ concentrations were detected after 1 week between 0.5–3 mM (Figure 3A). Most of the bioreactors maintained 1–4 mM headspace CO₂ throughout the remainder of the experiment. Some fluctuations were observed throughout the experiment but no consistent trends or significant differences were observed between the Cr levels. By 15 weeks, CO₂ concentrations in all bioreactors ranged between 1.0–2.7 mM CO₂. CH₄ was detected after 3 weeks and maintained a fairly steady concentration ranging between 0.1–1 mM. Substantially higher concentrations were observed in Lactate and Low-Cr bioreactors (0.5–2 mM) compared to High-Cr (0.01–0.15 mM CH₄).

Figure 2. Averaged metabolite concentrations in duplicate bioreactors inoculated with Hanford well H-100 groundwater. (a) Lactate and (b) Acetate concentrations when amended with (X) 0 mg/L Cr(VI), (○) 0.1 mg/L Cr(VI) and (▼) 3.0 mg/L Cr(VI). W = weeks. Error bars indicate 1 standard deviation between replicates.
doi:10.1371/journal.pone.0083909.g002
Cr(VI)-reduction

Immediate reduction of Cr(VI) was observed in Low and High-Cr bioreactors as Cr(VI) was below detection from week 1 through the duration of the experiment (data not shown). Abiotic reduction of Cr(VI) did not occur in the sterile media (data not shown). Washed cells from all bioreactors displayed the ability to reduce Fe(III) and Cr(VI), but in differing amounts throughout the experiment (Figure 4). After 4 weeks, the Cr(VI)-reduction rate in Cr amended bioreactors appeared to be higher than in the lactate only. Washed cells from lactate only bioreactors were able to reduce about 32 μM Cr(VI)/hr, whereas the Low-Cr and High-Cr bioreactor cells were able to reduce about 39 and 47 μM Cr(VI)/hr, respectively. However, by 7 weeks, the Cr(VI)-reduction rates significantly dropped to ~6–8 μM/hr in all the bioreactors, and subsequently remained in the range of 4–8 μM/hr. Slightly higher Cr(VI)-reduction rates were observed in High-Cr compared to other bioreactors. Opposite temporal trends were observed for Fe(III)-reduction rates. Between 4 and 8 weeks, the Fe(III)-reduction rates increased from 450 μM/hr to about 1000 μM/hr in lactate only bioreactors, from 550 to 820 μM/hr in Low-Cr and from 675 to 3050 μM/hr in High-Cr bioreactors. After 9 weeks, the Fe(III)-reduction rates decreased in all bioreactors, to 600, 1200 and 2100 μM/hr at 14 weeks in lactate, Low-Cr and High-Cr bioreactors, respectively. Washed cells from High-Cr bioreactors exhibited consistently higher Fe(III) reduction rates than...
Low-Cr and lactate bioreactors throughout the experiment except at 13 weeks.

**Microbial community composition**

Native microbial community composition from Hanford H-100 ground water significantly changed with lactate enrichment in the continuous flow bioreactors, including the ones amended with Cr(VI). NMDS analysis of Bray-Curtis distances of the rarefied Bacteria and Archaea OTUs between samples showed that major separation occurred between native and lactate amended samples (Figure 5). Microbial communities did not diverge much between Cr amended versus the lactate only controls. Temporal separations were evident initially, as bacterial communities from 3d and 1 week time points clustered together. However, after 1 week there was not much temporal separation. Similarly the Archaea OTUs at 3d, 1 and 2 weeks did separate from rest of the time points. After two weeks most samples between 10 and 13 weeks clustered together, but no distinct trends were observed between Cr treatments. Lactate, acetate and CO$_2$ correlated with bacterial OTUs separation among the metabolites tested. Archaea OTUs separation correlated mostly with lactate, acetate, CH$_4$ and H$_2$.

Relative proportions of the predominant bacterial taxa (Figure 6) also showed that native Hanford ground water communities changed significantly with lactate enrichment. Native bacterial communities were more diverse (Shannon diversity = 4.50 (Table S1 in file S1)), with higher relative proportions of *Tsproemonia* (a Spirochaete), *Thermodesulfobium* (a Firmicute), *Desulfovomonile* (a Deltaproteobacteria) and *Persivina* (a Bacteroidetes) (Figure 6). *Firmicutes* (18%), *Proteobacteria* (17%) and *Spirochaetes* (13%) were the predominant phyla detected in native environments (Figure S1 in file S1). After 3 days of lactate enrichment, 95% of the communities were *Sporotalea* spp., *Pelosinus* spp. and *Pseudomonas* spp. Among the native communities, they were detected at ~1.7, 1.4 and 2.1% of the total communities, respectively (Figure S1 in file S1). After 1–4 weeks in the lactate only bioreactor, *Sporotalea* spp. increased to ~70% of the total community and *Pelosinus* spp. to ~25–30% of the community. Species diversity within *Veillonellaceae* family was slightly different initially between Cr levels but clustered together after 5 weeks (Figure S2 in file S1). Relative proportions of *Pseudomonas* spp. decreased to <5%.

Similar trends were observed in low and High-Cr bioreactors. After 4 weeks the relative proportions of other bacteria such as *Brevundimonas* spp., *Sulfurospirillum* spp., *Rhizobium* spp., *Robinsoniella* spp. and a few others slightly increased with some temporal variations. These trends were also noticed in Low-Cr bioreactors after 2 weeks and in High-Cr bioreactors after 3 weeks. The diversity indices analysis indicated that the overall diversity of the communities increased at 3–4 weeks. There were no consistently significant differences in communities between Cr amended and the Lactate bioreactors.

The most abundance native Archaea genera in Hanford groundwater sample were *Methanoseta* (~55%), *Methanosarcina* (17%), *Halobacteriales* (12%), *Methanoregula* (8%) and others (Figure 7). After lactate enrichment, the Archaeal community composition significantly changed compared to the native community. By 3 days, most of the Archaea were unclassified groups (90%) and the rest were primarily *Methanosarcina*. None of the other predominant Archaea detected in native samples were prevalent after lactate enrichment. By 5 weeks, most of the Archaea classified as *Methanosarcina* (~95%) and remained until the end of the experiment. *Methanosarcina* barkeri was the closest relative for most of the predominant OTUs observed in bioreactors (Figure S3 in file S1). Similar trends were observed in Cr amended bioreactors. The number of OTUs detected and Shannon diversity index significantly decreased in all reactors compared to native communities (Table S2 in file S1). Archaea diversity continued to decrease over time and was significantly reduced after 5 weeks in lactate only and High-Cr reactors and after 3 weeks in Low-Cr reactors. However, High-Cr reactors established...
the opposite trend after 10 weeks with significantly higher diversity compared to others.

Microarray hybridization with GeoChip detected several groups of key functional genes including metal resistance, carbon, nitrogen and sulfur cycling genes. NMDS analysis of Bray-Curtis distances of normalized signal intensity of all the functional genes and key groups of genes are presented in Figure 8. Overall the functional gene composition among the bioreactors showed a greater separation of the High-Cr and Low-Cr communities, with the lactate only communities being more similar to High-Cr reactors, suggesting that the low level Cr exposure impacted the community populations more so than the higher level Cr insult. Separation of the communities between conditions correlated well with CO₂, acetate and butyrate production and lactate

Figure 5. Non-metric multidimensional scaling of OTU composition based on 16S rRNA gene sequences in bioreactors inoculated with Hanford well H-100 groundwater. (a) Bacteria and (b) Archaea amended with (X) 0 mg/L Cr(VI), (○) 0.1 mg/L Cr(VI) and (▽) 3.0 mg/L Cr(VI). W = weeks. doi:10.1371/journal.pone.0083909.g005

Figure 6. Temporal changes in the relative abundance of the most abundant Bacterial taxa detected by 16S rRNA gene sequencing in bioreactors inoculated with Hanford well H-100 groundwater. Bioreactors were amended with 0 mg/L Cr(VI) (Lactate), 0.1 mg/L Cr(VI) (Low-Cr) and 3.0 mg/L Cr(VI) (High-Cr). W = weeks. doi:10.1371/journal.pone.0083909.g006
Discussion

The results of this study indicate that a lactate enriched native ground water community from Hanford, WA reduced Cr(VI) and only some Archaea populations were impacted by Cr(VI) concentrations of 0.1 and 3.0 mg/L compared to lactate only treatment. Pelosinus spp. and Sporotalea spp., both from the family Veillonellaceae, were the predominant bacterial groups in all the bioreactors similar to our previous experiment [11]. Cr(VI) concentrations (3.0 and 0.1 mg/L) appeared to have not impacted the growth of these two groups of bacteria, which suggests that some of these species might have evolved Cr resistant strategies [11,12], and this may be as a result of long term Cr exposure at the Hanford sites. Possibly due to their Cr(VI) resistance, Pelosinus spp., have been observed to be one of the predominant groups of microbes detected from several sites with a history of Cr [10,12,30] as well as other metal contaminated subsurface environments [31]. It is also interesting that these bacteria were able to outcompete most other primary fermenters, sulfate and iron -reducers which may make these species ideal candidates for bioremediation of Cr impacted subsurface environments, particularly in environments limited by electron-acceptors such as sulfate.

Although overall Pelosinus spp. abundance did not diminish throughout the duration of the experiment, the Cr reduction potential of the community significantly decreased in all bioreactors after 4 weeks, which coincided with a slight increase in diversity. Other groups of bacteria such as Burkholderia, Comamonas testosteroni, Ralstonia pickettii seem to have proliferated more in Cr amended bioreactors. Other predominant Cr resistant bacteria detected included Burkholderia phymatum, Bradyrhizobium spp., Delftia acidovorans, Ralstonia solanacearum, and Shewanella spp. Non-metric multidimensional scaling of normalized signal intensity for the mcrA gene probes showed no consistent separation between Cr levels and lactate only bioreactors (Figure S5 in file S1). Some of the predominant Cr resistant bacteria such as Burkholderia vietnamiensis, Comamonas testosteroni, Ralstonia pickettii seem to have proliferated more in Cr amended bioreactors. Other predominant Cr resistant bacteria detected included Burkholderia phymatum, Bradyrhizobium spp., Delftia acidovorans, Ralstonia solanacearum, and Shewanella spp. Non-metric multidimensional scaling of normalized signal intensity for the mcrA gene (the alpha subunit of methyl coenzyme M reductase [29]) detected by GeoChip showed clear separation between Cr levels (Figure S4 in File S1).

Figure 7. Temporal changes in the relative abundance of the most abundant Archaea detected by 16S rRNA gene sequencing in bioreactors inoculated with Hanford well H-100 groundwater. Bioreactors were amended with 0 mg/L Cr(VI) (Lactate), 0.1 mg/L Cr(VI) (Low-Cr) and 3.0 mg/L Cr(VI) (High-Cr). W = weeks.

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consumption. Similar trends were observed for carbon, nitrogen and sulfur cycling genes, showing that Low-Cr communities were different from High-Cr and Lactate communities. Considerable numbers of chromium resistant bacteria (about 571 positive probes for chdA gene (a chromium efflux transporter [28]) were detected in these bioreactors. Hierarchical clustering of the normalized signal intensity of the chdA gene probes showed no consistent separation between Cr levels and lactate only bioreactors (Figure S5 in file S1). Some of the predominant Cr resistant bacteria such as Burkholderia vietnamiensis, Comamonas testosteroni, Ralstonia pickettii seem to have proliferated more in Cr amended bioreactors. Other predominant Cr resistant bacteria detected included Burkholderia phymatum, Bradyrhizobium spp., Delftia acidovorans, Ralstonia solanacearum, and Shewanella spp. Non-metric multidimensional scaling of normalized signal intensity for the mcrA gene (the alpha subunit of methyl coenzyme M reductase [29]) detected by GeoChip showed clear separation between Cr levels (Figure S4 in File S1).
The Cr(VI) concentrations were negligible in both Low and High-Cr reactors for the duration of the experiment, even though washed cell assays indicated that Cr(VI)-reduction potential decreased significantly after 4 weeks. These trends open up the possibility that some indirect Cr(VI)-reduction may have occurred via cell metabolites of the fermentative community along with expressed enzymes. A recent study suggested that Cr(VI)-reduction by a Pelosinus sp. was catalyzed by indirect mechanisms, as several cellular metabolites such as flavo-proteins were able to reduce Cr(VI) [12], which may have formed some Cr(III)-organic complexes as observed in other studies [35]. Some cellular toxicity to Cr(III) may have also occurred through transient Cr(III) ions [36] or Cr(III)-complexes which can be soluble and toxic [37], thereby possibly explaining the lower methane production observed in high-Cr reactors. Some of the Cr(III) may also be tied up with methanogenic substrates such as acetate, although it appears that other acetotrophic microbes were not impacted as acetate was still metabolized in High-Cr reactors. It seems that the methanogens may be particularly sensitive to Cr(III), an effect that should also be further investigated given the possibly important consequences to bioremediation. It is also critical to evaluate the long term stability of Cr-organic complexes toxicity and reoxidation to Cr(VI), although the latter seems to be unlikely based upon results from this study. Direct enzymatic reduction of Cr(VI) via

Figure 8. Non-metric multidimensional scaling of Geochip based functional genes composition. (a) all genes detected (b) carbon cycling (c) nitrogen cycling and (d) sulfur cycling genes detected in bioreactors inoculated with Hanford well H-100 groundwater amended with (X) 0 mg/L Cr(VI), (○) 0.1 mg/L Cr(VI) and (▼) 3.0 mg/L Cr(VI). W = weeks. doi:10.1371/journal.pone.0083909.g008
hydrogenases and cytochromes observed in several microorganisms [38] cannot be completely ruled out as *Pelosinus* spp. contain several hydrogenases and cytochromes [12]. Indirect Cr(VI) reduction by several microbial metabolites is also well documented. For example, under anoxic conditions, Cr(VI) can be readily reduced to Cr(III) by ferrous iron, sulfides and other low redox metabolites [4,39]. Perhaps a combination of these factors contributed towards non-specific Cr(VI) reduction and alleviated toxicity.

**Conclusions**

The results of this study showed that lactate enriched native ground water bacterial community from Hanford, WA did not significantly change with exposure to Cr(VI). Stable community structure was maintained with *Pelosinus* spp. and *Sporotalea* spp. and *Methanoarcana* spp. as the predominant groups detected at all three Cr(VI) levels, although CH4 production was significantly lower in High-Gr reactors. Cr(VI) was readily reduced in both Low and High-Gr bioreactors, mostly as a result of non-specific reduction by cellular metabolites. Results of this study suggest that low redox conditions with a stable fermentative community stimulated with an electron donor can be a viable option to keep Cr(VI) reduced over the long-term in environments lacking a sufficient supply of terminal electron acceptors.

**Supporting Information**

**File S1** Contains: File S1. Native microbial communities that were detected in Hanford well H-100 groundwater. (a) bacterial phyla and (b) Archaea classes. Figure S2. Non-metric multidimensional scaling of OTUs from *Veillonellaceae* family in bioreactors inoculated with Hanford well H-100 groundwater. Bioreactors were amended with (X) 0 mg/L Na2Cr2O7, (○) 0.1 mg/L Na2Cr2O7 and (▼) 3.0 mg/L Na2Cr2O7. W = weeks. Figure S3. Phylogenetic analysis of microbial community functional structures to pilot-scale uranium in situ bioremediation. Immune Journal 2: 177–183.

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