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The *Chthonomonas calidirosea* Genome Is Highly Conserved across Geographic Locations and Distinct Chemical and Microbial Environments in New Zealand’s Taupo Volcanic Zone

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ABSTRACT

*Chthonomonas calidirosea* T49 T is a low-abundance, carbohydrate-scavenging, and thermophilic soil bacterium with a seemingly disorganized genome. We hypothesized that the *C. calidirosea* genome would be highly responsive to local selection pressure, resulting in the divergence of its genomic content, genome organization, and carbohydrate utilization phenotype across environments. We tested this hypothesis by sequencing the genomes of four *C. calidirosea* isolates obtained from four separate geothermal fields in the Taupo Volcanic Zone, New Zealand. For each isolation site, we measured physicochemical attributes and defined the associated microbial community by 16S rRNA gene sequencing. Despite their ecological and geographical isolation, the genome sequences showed low divergence (maximum, 1.17%). Isolate-specific variations included single-nucleotide polymorphisms (SNPs), restriction-modification systems, and mobile elements but few major deletions and no major rearrangements. The 50-fold variation in *C. calidirosea* relative abundance among the four sites correlated with site environmental characteristics but not with differences in genomic content. Conversely, the carbohydrate utilization profiles of the *C. calidirosea* isolates corresponded to the inferred isolate phylogenies, which only partially paralleled the geographical relationships among the sample sites. Genomic sequence conservation does not entirely parallel geographic distance, suggesting that stochastic dispersal and localized extinction, which allow for rapid population homogenization with little restriction by geographical barriers, are possible mechanisms of *C. calidirosea* distribution. This dispersal and extinction mechanism is likely not limited to *C. calidirosea* but may shape the populations and genomes of many other low-abundance free-living taxa.

IMPORTANCE

This study compares the genomic sequence variations and metabolisms of four strains of *Chthonomonas calidirosea*, a rare thermophilic bacterium from the phylum *Armatimonadetes*. It additionally compares the microbial communities and chemistry of each of the geographically distinct sites from which the four *C. calidirosea* strains were isolated. *C. calidirosea* was previously reported to possess a highly disorganized genome, but it was unclear whether this reflected rapid evolution. Here, we show that each isolation site has a distinct chemistry and microbial community, but despite this, the *C. calidirosea* genome is highly conserved across all isolation sites. Furthermore, genomic sequence differences only partially paralleled geographic distance, suggesting that *C. calidirosea* genotypes are not primarily determined by adaptive evolution. Instead, the presence of *C. calidirosea* may be driven by stochastic dispersal and localized extinction. This ecological mechanism may apply to many other low-abundance taxa.

To date, the poorly characterized phylum *Armatimonadetes* is described primarily by environmental 16S rRNA marker gene data (1–4) and has only three described type species (5–7), from which two genomes have been sequenced (8, 9). Analysis of these genome sequences shows that *Armatimonadetes* are most closely related to the candidate lineages FBP and WS1 and to the phylum *Chloroflexi*. *Armatimonadetes* has three described classes, each represented by a single type strain: *Armatimonadia* (*Armatimonas rosea* YO-36 T) (6), *Fimbriimonadina* (*Fimbriimonas ginsengisoli* Gsoil 348 T) (7), and *Chthonomonadina* (*Chthonomonas calidirosea* T49 T) (5). The phylum contains at least eight additional class-level phylogenetic lineages without cultivated representatives (3).

*Chthonomonas calidirosea* T49 T is an aerobic and moderately acidophilic thermophile isolated from geothermal soil within the Taupo Volcanic Zone (TVZ), New Zealand (10), an area rich in geothermal systems and surface hydrothermal features. In addition to extracellular polymeric substances, the strain produces extracellular saccharolytic enzymes, which allow it to utilize many
diverse carbohydrates, with the exception of crystalline insoluble polymers (e.g., cellulose) (5). Genomic analysis of strain T49T identified a wide range of glycosyl hydrolases and carbohydrate ATP-binding cassette transporters, as well as many extracellular function sigma factors (8). Based on genomic sequence and physiological characterization, the ecological role of Chthonomonas calidirosea T49T was proposed to be that of a scavenger, utilizing heterogeneous carbohydrates from degraded biomass within the environment (5, 8). Notably, no Ch. calidirosea-like phylotypes have been reported from outside the TVZ (3, 11, 12). The most similar phylotypes (based on pairwise comparison, GenBank accession numbers KM102610 and KM102602 have 93% 16S rRNA gene sequence identity to strain T49T) originated from volcanic soils at Paricutin Volcano, Mexico (see also the supplemental material).

One remarkable feature of strain T49T is its apparent genomic disorganization. Many functionally related genes that are typically organized into operons in other bacterial genomes (e.g., histidine, tryptophan, and purine biosynthesis) are instead individually distributed throughout the C. calidirosea genome. This genomic disorganization complicates metabolic pathway predictions, although the abundant sigma factors observed in the genome provide a potential mechanism for gene regulation (8, 13). The high dispersal of functionally related genes observed in C. calidirosea appears to be uncommon. High genomic disorganization is observed in some cyanobacterial species (14–16), as well as in the Armatimonadetes species F. ginsengisoli (1) (see Fig. S1 in the supplemental material). Among the cyanobacteria, Prochlorococcus genomes are highly plastic and frequently reorganized by bacteriophages in response to their environment (17). Prochlorococcus genomes often display shorter and more dispersed operons than are found in other bacterial genomes (14, 18, 19). We therefore hypothesized that the nonoperonic genome organization of strain T49T would likewise be highly plastic and responsive to the selection pressure of the immediate environment, and this plasticity would be phenotypically reflected in traits such as the repertoire of carbohydrate utilization in distinct environments.

In order to assess the degree of genomic flux within the C. calidirosea genome in response to environmental selection pressure, we compared the genome sequences of strain T49T to those of three additional C. calidirosea isolates cultured from geographically isolated sites, averaging 46.2 km from the site of strain T49T isolation. In addition, we collected environmental geochemistry data and defined the microbial community structures by assessing the 16S rRNA gene sequence diversity for each sample site. With these data, we evaluated the relationships among geographic distance, geochemistry, community diversity, and genomic sequence variation in C. calidirosea.

MATERIALS AND METHODS

Cultivation of Chthonomonas calidirosea isolates. Based on 16S rRNA gene clone libraries constructed from diverse sites within the TVZ (data not shown), four geographically separated geothermal systems (Tikitere [TKT], November 2006; Waikite [WKT], July 2006; Wairakei [WRG], April 2009; and Te Kopia [TKA], September 2010) containing Armatimonadetes/candidate division OP10 were chosen for this study. Four pure cultures of C. calidirosea (T49T, P488, WRG1.2, and TKA4.10) were isolated from these sites under cultivation conditions previously described for oligotrophic carbohydrate-utilizing thermophiles (10) (Fig. 1). Strain T49T was physiologically characterized in a previous study (5).

Genome sequencing, assembly, and quality assessment. (i) Sequencing of C. calidirosea isolate genomes. After cultivation, genomic DNA was extracted from isolates P488, WRG1.2, and TKA4.10 using a previously described phenol-chloroform method (8). The four genomes were sequenced via different technologies (Table 1). Strain T49T was sequenced in a previous study (8). For isolates P488 and WRG1.2, genomic DNA
libraries were constructed using the Nextera XT DNA sample preparation kit (Illumina, Inc., San Diego, CA, USA), according to the manufacturer’s protocol, and 2 * 150-bp paired-end sequencing was performed using a 300-cycle sequencing kit (version 1.0) on an Illumina MiSeq. For isolate TKA4.10, a genomic DNA library was constructed using the Ion Xpress Plus fragment library kit with 100 ng of input DNA. The beads were prepared using the Ion One Touch 200-bp version 2 DL kit, and sequencing was performed with the Ion PGM 300-bp sequencing kit on an Ion Torrent (Thermo Fisher Scientific, Inc., Waltham, MA, USA).

(ii) Assembly and annotation of isolate genomes. MIRA version 4.0rc3 (20) was used to assemble the genomes of P488, WRG1.2, and TKA4.10. The strain T49† genome (8) was used as a reference during assembly. This resulted in a single scaffold for each genome, in which ambiguous regions were indicated with repeating Ns (IUPAC notation). The resulting assemblies were aligned using progressiveMauve (21). Assembly coverage (Table 1) was calculated from contigs of ≥5,000 bp to avoid inflation of the figures. Gene prediction and annotation of the genomes were performed using the Integrated Microbial Genome-Expert Review (IMG-ER) pipeline (22). Further details regarding sequencing, assembly, and quality control of genome assemblies are described in Fig. S2 and S3 in the supplemental material.

Genome comparison and phylogenetic analysis. (i) Identification of gene homologs across C. calidirosea isolates. To determine whether annotated genes were conserved between strains, we calculated the read coverage for each gene in each strain using Bowtie2 (23) and BEDTools (24). The complete code for calculating gene coverage is available at http://bitbucket.org/biobakery/chthonomonas. The identification of conserved homologs between genomes with UCLUST (25) and the identification of putative genes in unmapped reads are described in the supplemental material and are summarized in Fig. S4.

(ii) Determination of isolate phylogeny. The PhyloPhlAn gene collection of 400 ubiquitous and phylogenetically informative genes (26) was used for phylogenetic analysis of the isolates. The C. calidirosea genomes contained 327 of these genes (see Table S1 in the supplemental material). Due to the high similarity between the isolates, we conducted Bayesian selection of 400 ubiquitous and phylogenetically informative genes (26) using MrBayes (27). The genes were divided into partitions and were unlinked so that each partition had its own set of parameters for estimation during the Markov chain Monte Carlo (MCMC) process. Each partition used the general time-reversible substitution model, with the heterogeneity rate determined by the invariable site plus gamma distribution (GTR + I + G). The model was selected using jModelTest (29) with the Akaike information criterion (AIC). Two MCMC runs were conducted. Each MCMC process ran for 140 million generations, sampling every 1,000 generations, with 35 million (25%) generations as burn-in. MCMC performance diagnostics were conducted via Tracer (http://tree.bio.ed.ac.uk/software/tracer/).

(iii) Genomic rearrangement comparison with other thermophiles. We compared the degree of genomic sequence rearrangement observed in C. calidirosea to that observed in other thermophilic species isolated from defined geographic locations. The genomic sequences of Sulfolobus islandicus (strains Y.N.15.51 and Y.G.57.14) and Thermus thermophilus (strains HB-8, HB-27, and JL-18) were downloaded from NCBI GenBank, and the genomic rearrangement of each species was assessed by aligning all of its strains (including those of the C. calidirosea isolates) using progressiveMauve (21). PhyloPhlAn and MUSCLE were used to select and align conserved genes within the genome sequence of each compared species (i.e., C. calidirosea, T. thermophilus, and S. islandicus). Sequence identities/dissimilarities were calculated from the concatenated aligned gene sequences with SIAS (http://imed.med.ucm.es.Tools/sias.html). Due to an incomplete assembly (>100 contigs), T. thermophilus strain ATCC33923 could not be analyzed for genomic sequence rearrangement, but the genome sequence was used for assessment of the divergence of conserved genes. Further details are described in the supplemental material and are summarized in Fig. S5.

Characterization of C. calidirosea isolate metabolisms with Biolog phenotype microarrays. The substrate utilization phenotypes of the four C. calidirosea isolates were compared using Biolog phenotype microarrays (Biolog, Inc., Hayward, CA, USA). All isolates were maintained using 4.5NZS solid medium (10). Colonies were collected from the medium after 1 week of incubation at 60°C and were inoculated 250 mL of 4.5NZS liquid medium (pH 5.5), and 3 g/liter maltose and 0.2 g/liter Casamino Acids were added in 500-ml containers under an air headspace. The liquid cultures were incubated at 60°C for ~40 h. Cells from 50 mL of culture were pelleted by centrifugation (5,190 × g for 15 min) and washed with sterile water three times prior to resuspension in a maltose-free 4.5NZS liquid medium at an optical density at 600 nm (OD<sub>600</sub>) of 0.6 to 0.8. A 100-μL aliquot of the resuspended culture and 5 μL (5 g/liter) of MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide] were added to each of the 96 wells on a Biolog PM1 plate (Biolog, Inc.). The plates were sealed and incubated in darkness at 60°C for ~23 h. After incubation, 900 μL of dimethyl sulfoxide was added to each well (30), and the OD<sub>490</sub> was recorded. Each isolate was tested twice, using a separate cultivation batch for each experimental replicate. Mean responses were calculated from the corresponding wells of each isolate, and tie-corrected Spearman rank-order correlations between isolates were calculated with the R function cor.test (31) (see Fig. S6 and Table S9 in the supplemental material).

Analysis of soil chemistry and mineral content. Soil temperature was measured on site using a Fluke 50S thermocouple. Sample pH was measured in the laboratory at 25°C by mixing 1 g of the sample in 10 mL of deionized water. Soil moisture content and major oxides were analyzed using X-ray fluorescence (XRF). Major oxides were detected by XRF with borate fusion of the samples. The total soil carbon content was measured.
by mixing weighed soil aliquots with 25 ml of 0.05 M K$_2$SO$_4$ and filtering to extract total soluble carbon. Total soluble carbon was then measured bycombusting the samples and measuring the CO$_2$ evolution using a multi N/C 3100 analyzer (Analytik Jena AG, Jena, Germany). The inorganic carbon content of the sample was measured separately by quantifying the CO$_2$ generated by the acidification of an aliquot of the original sample (32). Mineral content was determined by X-ray diffraction (XRD) (Philip X’Pert Pro with Co K$_x$ radiation source) with the software X’Pert HighScore (Spectris plc, England) and Siroquant (Sietronics Pty Ltd., Canberra, Australia) (see Fig. S7 in the supplemental material). Soil nitrate, nitrite, ammonia, and dissolved reactive phosphate levels were determined by flow injection analysis (FIA) (Hach Company, Loveland, CO, USA). Analytes were extracted by mixing 1 g of soil sample with 50 ml of 1 M KCl. The mixture was centrifuged, and the resulting supernatant was passed through a 0.2-µm-pore-size filter prior to analysis.

Community 16S rRNA gene-targeted sequencing and bioinformatic processing. Total DNA was then extracted from each soil sample. Prior to DNA extraction, each soil sample from each site was mixed with 500 µl of a 50-g/liter sterile skim milk solution (Becton, Dickinson and Company, NJ, USA) to reduce DNA binding to clay. DNA was then extracted with a NucleoSpin soil DNA kit (Macherey-Nagel GmbH & Co. KG, Düren, Germany) according to the manufacturer’s instructions. The V4 hyper-variable region of the 16S rRNA gene was amplified by PCR and sequenced using the Ion Torrent platform, according to standardized methods, as described in detail in the supplemental material. Quality control was performed with UPARSE (33), as described in detail in the supplemental material, resulting in a mean of ~20,700 sequences per sample. These sequences were clustered de novo into operational taxonomic units (OTUs), with a minimum identity of 97%, and the representative sequences were taxonomically assigned using the UCLUST (25) consensus taxonomy assigner in QIIME 1.8.0 (34). Greengenes release 13.8 (35) was used as the reference taxonomic database. The OTU table was rarefied to 17,941 sequences per sample to remove sample heterogeneity; rarefaction analysis indicated that the sequencing depth was sufficient to identify the majority of OTUs within each community (see Fig. S8 in the supplemental material). QIIME (34) was used for β-diversity calculations and downstream analyses; β-diversity was measured by Bray-Curtis dissimilarity and by weighted and unweighted UniFrac indices. A weighted-UniFrac hierarchical clustering tree was constructed using the unweighted pair group method using average linkages (UPGMA). Jackknife analysis with the UPGMA tree was conducted with 200 repetitions and 12,000 sequences per sample. PyNAST alignment (36) and FastTree (37) were used to generate phylogenetic trees.

Accession numbers. The genome sequence and assembly data of the *C. calidirosea* isolates were deposited at EMBL/GenBank with the following accession numbers: PRJEB1573 (strain T49$^T$), PRJEB4907 (isolate P488), PRJEB4936 (isolate WRG1.2), and PRJEB4937 (isolate TKA4.10). Community sequencing data were deposited under accession number PRJEB13454.

**RESULTS**

The aim of this research was to understand how the ecosystem shapes the evolution of the *C. calidirosea* genome and phenotype. To connect genome dynamics with environmental factors and observed phenotypes, we isolated four *C. calidirosea* strains from four geographically distinct geothermal soil environments. For each site, we determined the soil chemistry and microbial community composition by 16S rRNA gene sequence analysis. We sequenced each isolate genome and performed nucleic acid sequence divergence assessment, phylogenetic inference, and gene content and organization analysis. In addition, we characterized the metabolic profile of each isolate (Fig. 1).

**Sample sites show major differences in clay content and hydrothermal activity.** The four sample sites were distributed along the TVZ (see Fig. S9 in the supplemental material), with a maximum linear distance of 67.4 km and minimum of 12.0 km between sites. All sample sites were geothermally affected soils. Soil pH ranged from 3.5 to 4.8, and the temperature ranged from 46.9 to 59.9°C. We characterized the mineral compositions of the sample sites using semiquantitative XRD and XRF analyses (see Fig. S7 and Table S2 in the supplemental material). The soil minerals were primarily quartz, silica, and clay. Clay minerals are sensitive to hydrothermal alteration. As shown in Fig. S7, the clay types at sites TKA and TKT were almost entirely primary clay minerals (e.g., magnetite, biotite, pyroxene, and plagioclase), indicating that these ecosystems were either newly affected or weakly affected by hydrothermal activity. In contrast, sites WKT and WRG had mostly secondary clay minerals (e.g., alunite-kaolinite and montmorillonite), which are formed from the reaction of rock with acidic steam-heated water over an extended period of time (38–40) (Table 2). WKT had the highest clay content, followed by WRG and TKT, while TKA was mostly quartz (86% [wt/wt]) (see Fig. S7). WKT was also much richer in soluble organic carbon (SOC) (2.33 mg/g of soil) than the other sites (range, 1.06 to 1.59 mg/g of soil) (Table 2). This SOC concentration was comparable to that of ambient-temperature soil environments, such as the nutrient-rich crop field soil (6.1 mg/g of soil) (41) and nutrient-poor pine plantation forest soil (0.08 to 0.17 mg/g of soil) (42), indicating that organic carbon is available at the sample sites.

*C. calidirosea* is a low-abundance taxon in diverse communities dominated by *Crenarchaeota* and *Thaumarchaeota*. We determined the relative abundance of *C. calidirosea* and other taxa at each sample site by community 16S rRNA gene sequencing. We observed a total of 190 operational taxonomic units (OTUs) among the four sites (see Table S3 in the supplemental material). A single OTU representing the genus *Chthonomonas* (OTU_085, with 99% sequence identity to strain T49$^T$) was detected at very low relative abundance (ranging from 6 × 10$^{-5}$ [TKA] to 3 × 10$^{-3}$ [WKT]) in all four communities (see Table S4 in the supplemental material). Although the short read length and error rate of ion semiconductor sequencing (Ion Torrent) cannot support

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**TABLE 2 General physicochemistry of soil samples**

| Sample site (isolate) | Temp (°C) | pH  | Moisture (%) | SOC (mg/g of soil)$^a$ | Ammonia (mg/g of soil) | Nitrate (mg/g of soil) | Nitrite (mg/g of soil) | Dissolved reactive phosphorus (mg/g of soil)$^a$ |
|-----------------------|----------|-----|--------------|------------------------|------------------------|------------------------|------------------------|----------------------------------|
| TKT (T49$^T$)         | 52.5     | 4.3 | 28.0         | 1.47                   | 145.0                  | 2                      | 0.15                   | BDL                              |
| WKT (P488)            | 50.7     | 4.5 | 59.9         | 2.33                   | 105.0                  | 6                      | 0.35                   | BDL                              |
| WRG (WRG1.2)          | 46.9     | 4.8 | 37.1         | 1.06                   | 28.5                   | 2                      | 0.40                   | BDL                              |
| TKA (TKA4.10)         | 59.9     | 3.5 | 28.8         | 1.59                   | 23.5                   | 1                      | 1.50                   | BDL                              |

$^a$ SOC, soluble organic carbon.

$^b$ BDL, below detection limit.
high-resolution comparison, these data confirm the detection of *Chthonomonas* with limited phylogenetic diversity (within the 97% OTU clustering criterion). The *C. calidirosea* OTU was most abundant in the more nutrient-rich soil at WKT, which had higher soluble organic carbon content, nitrate concentration, and overall clay mineral content (Table 2; see also Fig. S7 and Tables S3 and S4 in the supplemental material). In contrast, *C. calidirosea* was less abundant in sites consisting mainly of fresh hydrothermal deposits lacking in soluble organic carbon but rich in quartz and amorphous silica, as represented by the SiO2 concentration (see Table S2 in the supplemental material). Clay and organic matter increase the buffering capacity of soil (43), which may facilitate the survival of *C. calidirosea*, a species with a very limited pH range for growth (5, 8).

We observed only 21 archaeal OTUs, but these OTUs comprised the majority of the community at all four sites (56% [WKT] to 75% [TKA]). Five archaeal OTUs were present at all sites. Nearly half of observed archaeal OTUs (10/21) were *Euryarchaeota*, all belonging to the thermophilic acidophilic class *Thermoplasmata* (44). However, these represented <5% of the total sequence reads from any community. In contrast, the *Crenarchaeota* and *Thaumarchaeota* comprised the majority of reads from each community (Fig. 2). The *Crenarchaeota*, which were mainly *Thermoprotei* (nine OTUs [39 to 58% of total sequence reads]) (45), dominated all communities except TKT. The deep-branching MBG-A group (45–48) (12 OTUs [13 to 28% of total sequence reads]) dominated the TKT community (57% of OTUs). In this study, the class *Thermoprotei* was predominantly represented by the lineage NYPFEA, which is also associated with geothermal features at Yellowstone National Park, USA (49). Unfortunately, the lack of cultivated representatives within these *Crenarchaeota* and *Thaumarchaeota* lineages impedes any attempt to infer their ecological roles.

The four communities were distinctly different at a finer taxonomic resolution. Only 25 of the 128 total bacterial OTUs were observed at all four sites, and none of these were highly abundant (see Table S3 in the supplemental material). These taxa included *Sulfo bacteriaceae*, *Rhodospirillales*, and *Acidimicrobiaceae*. Based on their affiliations to characterized type strains, we speculate that these bacteria are moderately acidophilic, chemolithotrophic, and aerobic, and they grow via mechanisms, such as iron and sulfur oxidation (*Sulfo bacteriaceae* [50]), iron oxidation (*Acidimicrobiaceae* [51]), or methanotrophy (“*Candidatus Methylocystis*” [52]). In contrast, many predominant OTUs were detected in only one or two sites. For example, an archaeal OTU from clade SK322 (OTU_0003) comprised 10.3% of the total reads from WKT but was not detected at TKT, while an *Acinetobacter* OTU (OTU_158) comprised 8.5% of the total reads from WKT but was undetected at the other sites (see Table S3). These observations indicate that although the four *C. calidirosea*-hosting geothermal systems were broadly similar (in pH, temperature, and hydrothermally affected clays), they did not support similar microbial community compositions. This suggests that other ecological or stochastic differences determine the abundance of *Chthonomonas* spp. within these ecosystems. *C. calidirosea* genomic content and organization are highly conserved between isolates. The genomes of *C. calidirosea* isolates P488, WRG1.2, and TKA4.10 were sequenced using the Illumina MiSeq or Ion Torrent platform (Table 1) and assembled using the previously sequenced T49F genome (8) as a template. MIRA (20) and SPAdes (53) produced very similar assemblies (see the supplemental material), indicating little bias due to assembler choice. The length and G+C content of the TKA4.10 genome were identical to those of T49F; this was potentially influenced by the inefficient assembly of single-ended Ion Torrent reads beyond reference contig boundaries. Isolates P488 and WRG1.2 had slightly longer genome assemblies and lower G+C content than T49F and TKA4.10 (Table 1).

We determined the phylogeny of the strains by aligning the concatenated nucleic acid sequences of 327 phylogenetically informative genes (26) (see Table S1 in the supplemental material) and building a phylogenetic tree using Bayesian inference (Fig. 3). The pairwise sequence identity of these genes was very high (average, 99.2%) between the isolates. Based on these data, strain T49F shares the highest similarity with TKA, followed by P488 and WRG1.2. The posterior probability support for this unrooted tree topology was >99%. Within this tree, TKA4.10 showed a very short branch length from the internal node shared with T49F. WRG1.2 had the longest terminal branch length. This is in slight disagreement with strain-level conservation of the 16S rRNA gene sequences; each genome had two copies of the 16S rRNA gene, and all are identical except in strain P488, in which the two copies differ by 1 base.

While the four *C. calidirosea* genome sequences showed a high degree of synteny in most genomic regions (see Fig. S2 and S5 in the supplemental material), a previous study identified two putative horizontally transferred regions, A (between the loci CCALI_00447 and CCALI_00449) and B (between the loci CCALI_00804 and CCALI_00807) (8) (see Fig. S10 in the supplemental material). These two regions were conserved between isolates T49F and TKA4.10 but are mostly absent in isolate WRG1.2. Isolate P488 contained only a small deletion in region B, and the sequences in both regions were slightly less conserved. The variability in these regions is consistent with our inferred strain phylogeny (Fig. 3); there is low divergence between T49F and TKA4.10, more divergence in P488 (e.g., deletions, truncations, and frameshift mutations in the loci CCALI_00449 and CCALI_00805), and the regions were either never acquired or deleted from WRG1.2. Finally, we compared our Bayesian-inferred strain phylogeny to the structure of the microbial community from each sample site, as measured by β-diversity (Fig. 3; see also Table S5 in the supplemental material). When communities were compared by metrics that consider OTU abundance (e.g., weighted UniFrac and Bray-Curtis dissimilarity), sites WRG and TKA had the most similar communities, followed by WKT and then TKT. In contrast, isolates TKA4.10 and T49F are phylogenetically the most similar, indicating that the differences in genome sequence phylogeny of *C. calidirosea* isolates are not reflected by differences in the community composition of their respective isolation sites.

Functionally annotated isolate-variant genes are primarily DNA-interacting enzymes. We next assessed the similarity of genetic contents between the genomic assemblies by aligning the reads from each isolate to the genes from all four isolates and then assessing the coverage of each gene in each strain (see the supplemental material). Most gene homologs (~2,600) were present in all four *C. calidirosea* isolates, but we identified 769 putative isolate-variant genes, many of which encoded hypothetical proteins or domains of unknown function (see Table S6 in the supplemental material). There were a total of 50 apparently isolate-unique
genes among the four isolates; the majority (31/50) of these were hypothetical proteins. Of the remainder, the most common functions were related to DNA modification (e.g., restriction enzymes and methylation).

Finally, we examined the genome sequences of the isolates for genes that were not assembled into scaffolds due to the use of strain T49T as a reference during assembly (i.e., genes absent from the T49T genome sequence but present in other isolates). We performed a de novo assembly of the unassembled reads for each genome and then used a blastn search (NCBI nr database) to search these contigs for genes. In this manner, we identified 14 additional putative functionally annotated genes absent from strain T49T (see Table S7 in the supplemental material).

The *C. calidirosea* genome is more highly conserved across geographic distance than *Sulfolobus islandicus* or *Thermus thermophilus*. The four *C. calidirosea* isolates exhibited very low

![Image of a microbial community tree](https://example.com/microbial-community-tree.png)

**FIG 2** Microbial communities of the sample sites. Each ring on the outside of the tree corresponds to a sample site, and the color intensity at each position on the ring indicates the relative abundance of the corresponding taxon at that sample site. The color intensity has been scaled nonlinearly to emphasize low-abundance taxa by transforming the relative abundance with the exponent (⅓). Cross symbols at the terminal nodes indicate representative reads with only higher-level taxon assignments and may contain multiple genera. The bacterial phylum Armatimonadetes is highlighted in yellow, and the member genus *Chthonomonas* of class Armatimonadetes (labeled “A”) is colored in magenta. The relative abundance of *Chthonomonas*-related OTUs at each of the four sample sites was as follows: WRG, 4.08 × 10⁻⁴; WKT, 3.02 × 10⁻³; TKA, 5.6 × 10⁻⁵; and TKT, 1.17 × 10⁻⁴.
genomic divergence across a substantial geographic distance (see Fig. S9 in the supplemental material). We compared the nucleic acid sequence divergence (0.33% to 1.17%) (Table 3) and genomic sequence rearrangement (see Fig. S5 in the supplemental material) of the C. calidirosea isolates with two other well-studied groups of thermophilic microorganisms: *Thermus thermophilus* and *Sulfolobus islandicus* (54, 55). The sets of both *T. thermophilus* and *S. islandicus* strains were isolated across a geographic scale comparable to that of the C. calidirosea cohort (see the supplemental material). Relative to C. calidirosea, nucleic acid sequence divergence was more than double (2.45%) in *S. islandicus* and more than triple (3.18% to 7.49%) in *T. thermophilus* (see Table S8 in the supplemental material). Both of these species also exhibited greater genomic sequence rearrangement across strains than was evident in any two *C. calidirosea* genome sequences (see Fig. S5).

Substrate utilization is conserved across *C. calidirosea* isolates, but the relative activities are isolate specific. Strain T49<sup>T</sup> was physiologically and genomically characterized as a scavenger capable of utilizing a wide range of carbohydrates (5, 8). To assess the extent to which each strain had physiologically adapted and specialized to its environment, we conducted carbohydrate utilization redox assays with Biolog PM1 carbon source microtiter plates. The plates contained 95 substrates, including mono- and disaccharides, amino acids, sugar alcohols, nucleosides, organic acids, and other potential energy sources. All isolates had similar substrate utilization profiles, with the highest redox response to hemicellulosic sugars (e.g., xylose and mannose) and negligible responses to amino acids as a sole carbon and energy source. However, the relative intensity of substrate response was isolate specific and reproducible. We assessed isolate substrate response similarity by calculating the Spearman rank-order correlation coefficients between the carbon utilization profiles (see Fig. S6 and Table S9 in the supplemental material). The most similar utilization profiles were from P488 and WRG1.2 (r = 0.943), while the least similar were T49<sup>T</sup> and WRG1.2 (r = 0.768). These substrate utilization differences do not appear to be due to variation in genes related to carbon metabolism. They may, therefore, be due to differences in regulatory genes or polymorphisms in functional genes, such as enzymes or transporters.

### DISCUSSION

**Role of *C. calidirosea* as a heterotrophic scavenger in microbial communities.** Our 16S rRNA gene analysis of the four *C. calidirosea*-associated communities revealed that *Crenarchaeota* and *Thaumarchaeota* dominated all communities, with diverse bacterial species as minor components. The variation in site community composition is likely influenced by their varied degrees of hydrothermal activity and different geochemistries. A previous culture-based study of the sample site WKT (10) resulted in the isolation of several bacterial taxa (including *C. calidirosea*) but no archaeal species. The dominant archaeal lineages detected in this study are only distantly related to any characterized isolates. How-
ever, based on the metabolic capabilities of described strains within related lineages (e.g., *Sulfolobales*, *Nitrosporaeales*, and *Nitrospumilales*), lithotrophic lifestyles would seem likely (56–59). We therefore speculate that the dominant crenarchaeotal and thaumarchaeal phylotypes may occupy the niche of autotrophic primary producers, supporting diverse, yet low-abundance, chemoheterotrophic bacterial species, like *C. calidirosea*. The scavenger phenotype of *C. calidirosea* may be well suited to persist in an ecosystem with minimal or inconsistent saccharide sources (5, 8). A recent study of bacterial communities in a Thailand hot spring also indicated a potential relationship between *Armatimonadetes* and autotrophs. This survey detected abundant *Armatimonadetes* and Chloroflexi OTUs associated with cyanobacterial mats (60). Pairwise contiguous megaBLAST analyses between the Thailand hot springs *Armatimonadetes* OTUs and *C. calidirosea* showed that the Thailand OTUs had low sequence similarities (75 to 84%) to the type strains of the three *Armatimonadetes* classes (see Table S10 in the supplemental material).

**Low genomic diversity in the face of geographical isolation.** Although the four *C. calidirosea* isolates were cultured from geographically distant sites across the TVZ, their genome sequences were highly conserved, particularly compared to other thermophilic microorganisms recovered across similar geographical scales (see Fig. S5 and Table S8 in the supplemental material). Phylogeny inferred from single-nucleotide polymorphisms (SNPs) and horizontal gene transfer indicated a close relationship between all isolates, particularly between T49T and TKA4.10. (SNPs) and horizontal gene transfer indicated a close relationship between all isolates, particularly between T49T and TKA4.10. Gene presence/absence analysis identified relatively few variant genes (see Tables S6 and S7 in the supplemental material), such as restriction-modification systems. These genes are known to be rapidly evolving (61) and mobile (62), which may explain their presence in an otherwise largely conserved pan-genome. Physiologically, all four *C. calidirosea* isolates had similar carbohydrate metabolism. Future comparative transcriptomic and proteomic analyses of the isolates may shed light on subtle underlying regulatory processes resulting in the variations of phenotypic response within these highly conserved genome sequences.

**Potential mechanisms underpinning genomic conservation across geographic distance.** The divergence between genome sequences of *C. calidirosea* isolates consisted primarily of SNPs, which may accumulate by genetic drift during geographic isolation. However, the phylogenetic distances of the isolate genome sequences do not reflect the geographic relationships of the sample sites (see Fig. S9 in the supplemental material). The reason for the low genomic sequence divergence between the *C. calidirosea* isolates despite their geographical and ecological isolation is not immediately clear. One possible explanation for the high genomic sequence conservation is a relatively recent sympatric determination of species of *C. calidirosea* from a mesophilic relative, leading to its occupation of a new niche in geothermal environments. A review of 16S rRNA gene data sets available in the NCBI and EMP databases indicates that *C. calidirosea* can presently be considered endemic to New Zealand, with no genus-level phylotypes detected elsewhere (see the supplemental material). A recent review of the environmental distribution of *Armatimonadetes* phylotypes indicated that temperate soil is the most dominant environment for the phylum (3). Of 39 phylotypes identified representing the class *Chthonomonadetes* (group 2), only two phylogenetically distant clones were associated with thermophilic environments. In contrast, group 10 (A and B) consists entirely of phylotypes from geographically disparate geothermal environments. Thus, the occasional occurrence of thermophily in the *Chthonomonadetes* may indicate either recently acquired thermophilic adaptation or sampling bias. The identification of more closely related phylotypes in the future will help clarify the evolutionary history and environmental distribution of *Chthonomononas*.

Although this hypothesis may explain the low diversity from an evolution and sampling perspective, it is inadequate to explain the very high similarity in the genome sequences from genetic drift, unless the dispersal of *C. calidirosea* was relatively recent. The high similarity between strain T49T and isolate TKA4.10 suggests recent dispersal between the two sites. A recent study found that thermophilic microbes can be globally dispersed via aeolian transport but are selected by their environments (63). In dynamic geothermal environments, taxa with very low abundance, such as *C. calidirosea*, may become locally extinct but subsequently recolonized from another site, resulting in high genetic similarity between two populations. Over a sufficiently rapid timespan of extinction and colonization, the genome sequences would reflect stochastic aeolian dispersal rather than adaptive responses to selection pressures of specific environments. In contrast, an abundant species would be less sensitive to the stochastic process of localized extinction and repeated seeding and thus more likely to present a clearer relationship between geography and phylogenetic distance. We believe the low diversity of *C. calidirosea* genome sequences potentially reflects these scenarios. While surface and groundwater water flow presents an alternative mechanism of dispersal between sites, this is unlikely due to the arrangement of river catchments and aquifers flowing away from individual sites (see Fig. S9). The environmental persistence of *C. calidirosea* is further restricted by its inability to sporulate and by its fastidious growth requirements (limited pH and temperature growth ranges). Thus, rapid dispersal between suitable habitats is a possible mechanism for maintenance of local populations and genome consistency.

Based on analysis of the genome sequences of *C. calidirosea* isolates, we reject our initial hypothesis that *C. calidirosea* genomic disorganization reflects plasticity and niche-specific adaptation. Instead, the genome sequence is highly conserved and shows a small degree of genetic drift rather than response to different ecological selection pressures at different sample sites. To date, *C. calidirosea* has been detected only within New Zealand’s TVZ, and its global distribution is currently unknown. If additional isolates can be cultured from greater geographic distances (such as the fumarolic/volcanic-associated phylotypes of Paricutín Volcano, Mexico, which share 93% 16S rRNA gene sequence identity), they will continue to improve our understanding of how the *C. calidirosea* genome is shaped by genetic drift. Rather than possessing a dynamic and highly plastic genome, the conservation of gene content and order (synteny) of *C. calidirosea* may be comparable to those of prokaryotes such as *Haloquadratum walsbyi* (64) and members of the SAR11 (*Pelagibacteriales*) clade (65), which are known for high genomic conservation.

**Conclusion.** In order to investigate the relationship between the *C. calidirosea* genome and its environment, we performed comparative genomics and determined the physiology of isolates obtained from four distinct sites. These data were integrated with microbial community analyses and environmental geochemistry data for the respective sites. We have shown that *C. calidirosea*, a thermophilic non-spore-forming bacterium, exhibits detectable,
albeit minor, genome sequence differences across a geographic distance. We propose that this bacterium is capable of dispersal across geographical barriers, and that the resulting genome conservation over space is potentially applicable to many low-abundance microbial taxa. Our approach shows the value of augmenting amplicon profiling of poorly characterized communities with additional data, such as the physiological characterization of isolates and geochemical analysis, in order to provide context and validation of ecological inferences.

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