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Metagenomic Bacterial Finshing at the JGI

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Metagenomic Bacterial Finishing at the JGI

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Desulforudis audaxviator

sections being continuous sequence.)

Korarchaeum cryptofilum OPF8

means. Its small size was helpful.

than with D. audaxviator.

Orchid (available from the Stanford Genome Center) at right

show clearly the low complexity of this community. (Orchid displays an ace file distributed around a circle, the heavy blue

From a more complex thermophilic community, it was

Though there was considerable enrichment, there is a great deal more representation of other species in this project

about 100 reads remain unincorporated out of 28000.

maintained as a community in the lab and enriched by

Accumulibacter phosphatis Type IIA str. CU-1

differential lysis. This organism was finished by conventional

These first two organisms certainly represent the

was presented by A. phosphatis. It was the first community

more than half and which eventually produced the assembly

finishing the genome. It allowed the bridging of many small

of the initial reads have fused into the final consensus There is moderate polymorphism. It was our good

Abstract: Metagenomic Bacterial Finishing at JGI.

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The sequencing to completion of uncultured bacterial genomes from mutualistic communities is a demanding process in the best of cases. The complexity of the community, the quantity of genomic DNA available, the fraction of the total DNA collected representing the organism under study are all added to the normal difficulties of establishing a

complete sequence. At the JGI we have managed to complete the sequence of three metagenomic organisms, and are investigating a fourth, presenting a considerable

range of difficulty.

Candidatus Korarchaeum cryptofilum OPF8, (NC_010482; GI:170289627), is the first of this apparently ancient hyperthermophilic phyletic group to be sequenced (3). The ability to obtain ample une applicativy answert preferencing provide group for the steadards (2). The annual for locating and the DNA of near-monoclural purposed how strain complexity made this the most straightforward sort of metagenomic subject. The target organism constitutes –40% of the Yellowstone thermal Obsidian Food community. The community could be maintained in culture, and it was solution that K. cryptoffilm was the most resistant member to SDS lysis, thus allowing DNA purification to better than 90%. Its strain complexity was low as indicated by a SNP rate of ~0.2%.

Some organisms have remote or difficult habitats limiting the availability of source material. This is the case with the thermophile Desulforudis audaxviator, (NC_010424; G1169830219), from fractures in the earth's crust at a depth of 2800 meters in a South African gold mine (4).

The bacteria were collected on filters through which large amounts of subterranean water wa

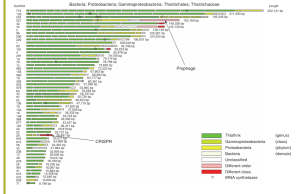
passed. The supersitis feed context on this aconystem contained but one species fortunately mean the but Networks and the the DNA yield of this one-time-only collection was sufficient to complete the genome. Considerably more complex situations are the rule, as illustrated by the case of *Candidatus Accumulibater phosphatis Type IIA str. CU-1*. This and closely related species are the principal actors in the Accommission of prosphilar stype LA six-Co-7. In a sing CoopSy Lanca spectra at the puncphil scions in use sequestation of morganic plosphilar as intracellular polyphosphate in wastewater treatment facilities. Bioreactor sludge encoded from a working facility in Wisconsin is a physically unresolvable mixture of organisms, which *Aphrophatis* producing that plus out 40%. The entite DAA sample was asquenced and the

resulting data thein subjected to phylogenetic graving using the Phylopythia (2) binning technique, greatly reducing the complexity of the subjected bin bin and the phylopythia (2) binning technique, greatly reducing the complexity of the subjecte bin bin and the phylopythia (2) binning technique, greatly reducing the complexity of the subject of the phylopythia (2) bin and the phylopythia (2) bin and the are now approaching a still more difficult genome. *Condidatis Ethodic robust relations* and the phylopythia (2) bin and the bulk plant materials as CO2-neutral fuel stocks.

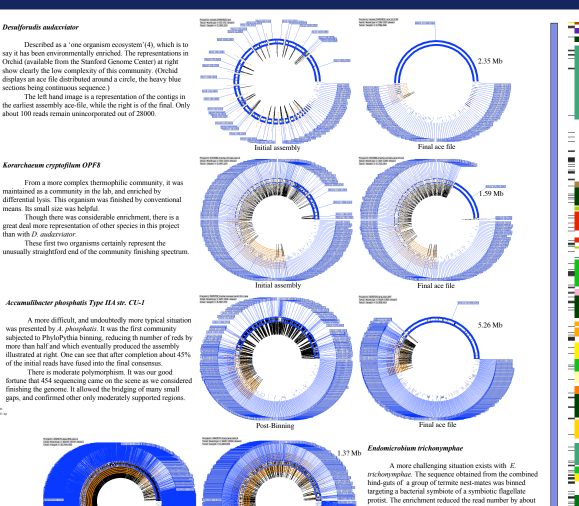
Again partitioning with Phylopythia was absolutely necessary. So far this has yielded some 42,000 Sanger reads. Of these about 40% of the assembled contigs are similar to related organisms. Also contig from 454 pyrosequencing contributed about 750,000 bp of additional coverage (-0.5x). Additional difficulty arises from sample size and strain complexity. There are several closely related organisms with substantial representation. There are several hindguts in the sample, and the lines of descent may be relatively independent even at the protist level.

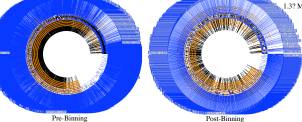
 Martin, H.G., Hugenholtz, P., et al. Nature Biotechnology, v24, no.10, 1263-69.
2006. Metagenomic Analysis of two Enhanced Biological Phosphorous Removal (EBPR) sludge communities (2) McHardy et al. Appl, Environ Microbiol 2000, 66(3):1175-1182, Accurate phylogenetic (a) Methady et al. Appl. Lawron Microbiol 2000, 00(5):1115-1162, Accurate p. c classification of variable-length DNA fragments.
(3) James G. Elkins, et al., PNAS. 2008 (in Press), The Korarchaeota: Archaeal

orphans representing an ancestral lineage of life (4) Dylan Chivian, et al., Science 2008 (in Press), Environmental genomics reveals a single species ecosystem deep within the Earth.



PhyloPythia binning of Thiothrix species (fron McHardy et al.(2) Scaffolds which consist of mate-pair linked contigs, 62% of which were binned by Phylopythia at the level of genus and 32% more at phylum level, with only 3% unclassified. Thiothricales constituted the largest clade of gammaproteobacteria in an Australian waste treatment sludge community studied for it's Accumulibacter species in parallel with the US species sequenced at the JGI.





organism project # enrich. mode degree of en GC Mb polym 454 subclone reads D. audaxviator 4000602 environ. 100% 62% 2.4 low Y 28167 K. crvptofilum 4001584 pchem 90% 49% 1.6 low Ν 19688 binning 3635729 40% 57% 5.3 mod 60579 A. phosphatis E. trichonymphae 4043170 binning < 40% 38% 1.3? high 37300 Y

trichonymphae. The sequence obtained from the combined hind-guts of a group of termite nest-mates was binned targeting a bacterial symbiote of a symbiotic flagellate protist. The enrichment reduced the read number by about 60%. The pertinent portion of the assembly at this point seems to consist of a number of strains considererably more distant from each other than the polymorphic members of the previous projects. This can be illustrated by the cartoon on the right.

In it, contigs with clear BLAST related sequence to a very closely related organism are aligned against that genome (AP009510.1), which is represented by the left vertical bar. The coloring is arbitrary to make contigs distinct.

There are certainly limits to the genomic resolution of communities, but luck and ingenuity have not been exhausted yet.

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