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Title

Single Cell Genome Reconstruction of Two Uncultured, Proteorhodopsin-containing Flavobacteria

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Abstract

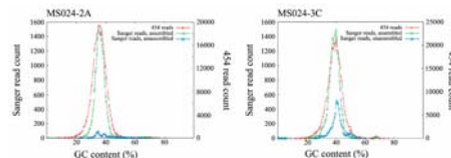
Determining the genetic makeup of predominant microbial taxa with specific metabolic capabilities remains one of the major challenges in microbial ecology and bioprospecting, due to the limitations of current cell culturing and metagenomic methods. The complexity of microbial communities and intraspecies variations hinders the assembly of individual genomes from metagenomic shotgun libraries. Here we report the use of single cell genomics to access the genome of two proteorhodopsin-encoding flavobacteria from Gulf of Maine bacterioplankton. We use high throughput fluorescence-activated sorting of single cells, whole genome amplification via multiple displacement amplification, PCR-screening and subsequent shotgun sequencing of these single amplified genomes (SAGs), allowing the genomic analysis of their novel photometabolic system and the sequence comparison to environmental marine sequence data.

Sequence assemblies

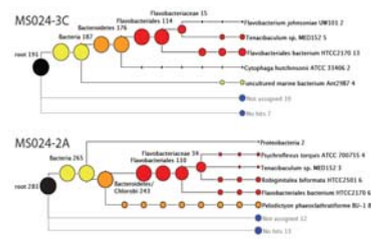
General features of the flavobacterial SAG assemblies.		
	MS024-2A	MS024-3C
Assembly statistics		
Assembly size [bp]	1,905,484	1,515,248
Estimated genome size [bp]	2,156,286 - 3,004,105	2,307,484 - 3,726,020
Number of contigs	17	21
Largest contig [bp]	684,032	549,383
GC content [%]	36	39
Mean read depth (± s.d.)	56 (± 63)	83 (± 110)
454 reads	47	68
Sanger reads	9	14.3
Gene predictions		
Total genes	1,824	1,426
Protein coding genes	1,785	1,400
with function prediction	1,205	960
w/o function prediction	580	440
Number of rRNA operons	2	1
Number of rRNA genes	33	24

The sequence data of the SAGs was Phrap assembled, followed by primer walking on shotgun clones, and PCR/adaptor PCR on the diluted MDA products.

Data QC

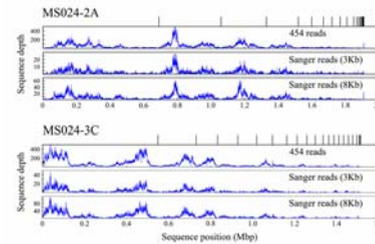


GC contents histogram of the unassembled and assembled Sanger and pyrosequencing reads for the two SAG exhibits a tight uni-modal distribution.

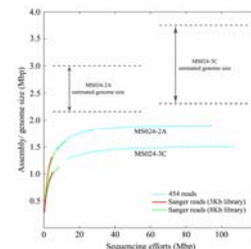


Taxonomic assignment analysis. The taxonomic contents of the blast output for the unassembled reads of the Flavobacteria sp. MS024-2A was estimated and visualized using the Metagenome Analyzer (MEGAN) (Huson, Genome Res 2007).

The MDA bias



MDA bias as evaluated by sequence depth distribution. The contigs for the SAG are aligned by length and contig breaks are indicated by the tick marks along the top. The mean sequence depth is 56 (± 63) for MS024-2A and 83 (±110) for MS024-3C.

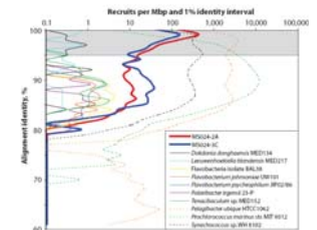


Genome coverage as function of the genome sequencing effort for the flavobacterial SAG. The curve displays near-saturation indicating that additional sequencing would mostly result in repeated sampling of the over-amplified genomic regions, not targeting the yet missing part of the genome.

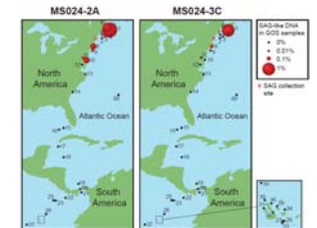
Chimeras

	MS024-2A		MS024-3C	
	chimeric reads/ clones (%)	overall chimerism	chimeric reads/ clones (%)	overall chimerism
Read-based chimerism				
30x Sanger reads (unsorted MDA DNA)	1.9	1 chimeric 28 Kbp	NA	NA
30x Sanger reads (5x treated MDA DNA)	2.0	1 chimeric 23 Kbp	1.9	1 chimeric 33 Kbp
30x Sanger reads (25 treated MDA DNA)	2.1	1 chimeric 22 Kbp	1.6	1 chimeric 40 Kbp
454 reads (5x treated MDA DNA)	1.8	1 chimeric 19 Kbp	1.6	1 chimeric 29 Kbp
454 reads (25 treated MDA DNA)	1.8	1 chimeric 21 Kbp	1.6	1 chimeric 29 Kbp
Clone-based chimerism				
30x Sanger (unsorted MDA DNA)	14.5	1 chimeric 23 Kbp	NA	NA
30x Sanger reads (5x treated MDA DNA)	5.2	0.2	0.2	0.2
30x Sanger reads (25 treated MDA DNA)	16.8	1 chimeric 15 Kbp	16.4	1 chimeric 20 Kbp
30x Sanger reads (unsorted MDA DNA)	14.5	1 chimeric 23 Kbp	NA	NA
30x Sanger reads (5x treated MDA DNA)	7.8	0.6	0.6	0.6
30x Sanger reads (25 treated MDA DNA)	3.0	0.3	0.3	0.3
30x Sanger reads (unsorted MDA DNA)	1.1	0.1	0.1	0.1
30x Sanger reads (5x treated MDA DNA)	1.0	0.1	0.1	0.1
30x Sanger reads (25 treated MDA DNA)	3.2	0.4	0.4	0.4
30x Sanger reads (unsorted MDA DNA)	36.4	1 chimeric 17 Kbp	28.5	1 chimeric 27 Kbp
30x Sanger reads (5x treated MDA DNA)	22.0	0.5	0.5	0.5
30x Sanger reads (25 treated MDA DNA)	2.5	0.2	0.2	0.2
30x Sanger reads (unsorted MDA DNA)	7.4	0.4	0.4	0.4
30x Sanger reads (5x treated MDA DNA)	3.3	0.3	0.3	0.3
30x Sanger reads (25 treated MDA DNA)	1.0	0.1	0.1	0.1
Average (all clones)		1 chimeric 17 Kbp		1 chimeric 23 Kbp

Fragment recruitment



Global Ocean Sampling (GOS) (Rusch, PLoS Biol 2007) metagenome fragment recruitment by the SAGs MS024-2A and MS024-3C, the currently sequenced marine Flavobacteria isolate genomes, the non-marine *F. johnsoniae*, and the three best GOS fragment recruiters *Pelagibacter*, *Prochlorococcus* and *Synechococcus*.



Geographic distribution of the GOS (Rusch, PLoS Biol 2007) metagenome fragments with >95% nucleotide identity to MS024-2A and MS024-3C.

Conclusion

Using the single cell approach, we demonstrate how a combination of single cell FACS and amplification via MDA can be used to access the genomes of uncultured environmental microorganisms, representative of their given environment.

Acknowledgements

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Background & Methods

