

1. Project Information

Program	Microbial
JGI Sequencing Project ID	1063230
Sequencing Project Name	Nostoc sp. Hyl-06-6-4

2. Read Statistics

	Raw Reads	Filtered SubReads	Error Corrected Reads
Reads	1,001,901	330,030	12,711
Bases	3,170,437,417	1,221,170,935	89,789,703
Avg Read Length	3,164.4 +/- 3,542.6	3,700.2 +/- 2,442.3	7,063.9 +/- 4,264.5
Reads >5 kbp	205,620	66,739	8,197
Bases, reads >5 kbp	1,797,115,030	505,461,320	79,902,869
Avg Read Length, reads >5 kbp	8,740.0 +/- 3,672.6	7,573.7 +/- 2,488.8	9,747.8 +/- 2,604.1

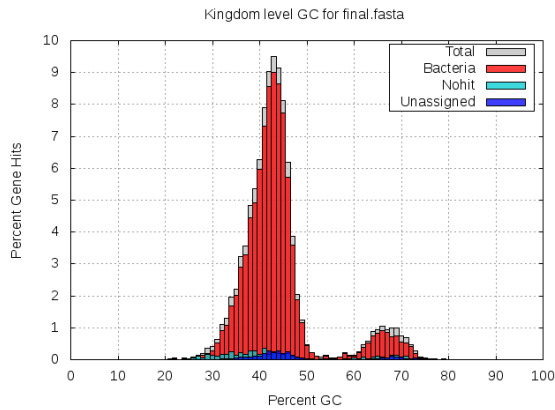
3. Assembly Statistics

Scaffold total	178
Contig total	178
Scaffold sequence length	7.628 mb
Contig sequence length	7.628 mb 0.000% gap
Scaffold N/L50	1/6.184 mb
Largest Contig	6,183.9 kbp
Number of scaffolds >50 kb	7
Pct of genome in scaffolds >50 kb	91.86%

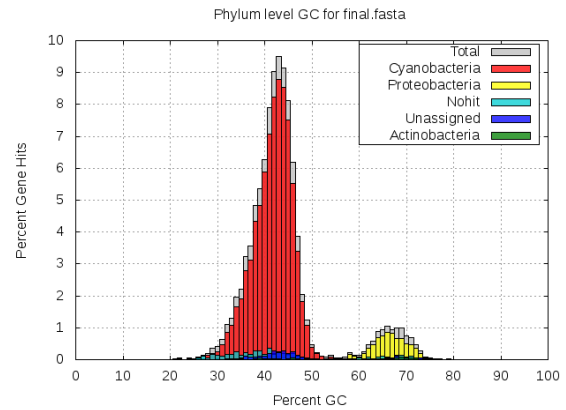
4. Assembly QC Results

GC histogram of the predicted genes on each contig, overlaid with GC of hits based on LAST, shown for different taxonomic levels.

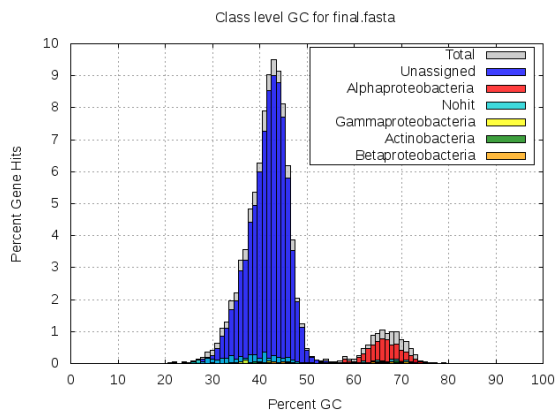
Kingdom



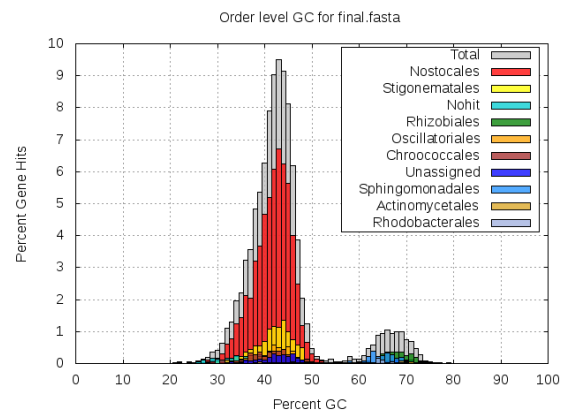
Phylum



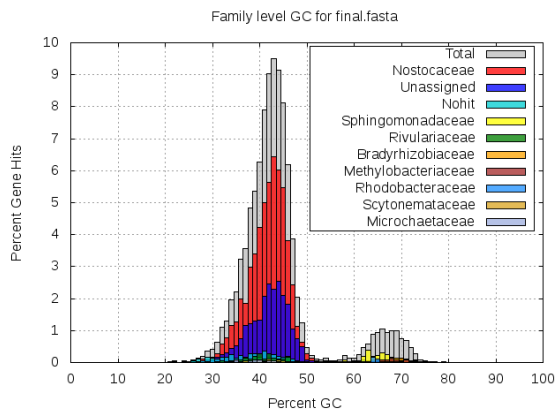
Class



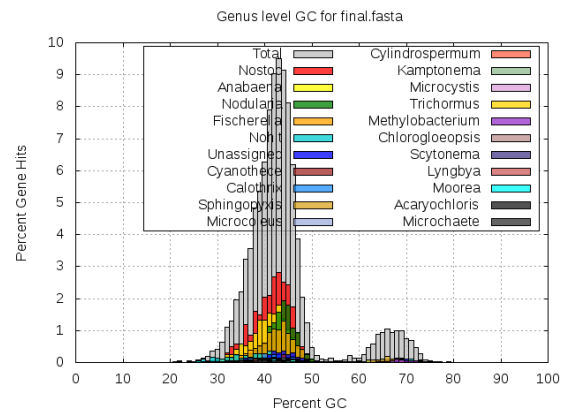
Order



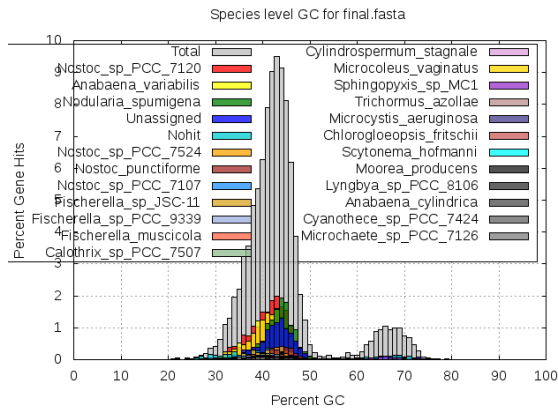
Family



Genus

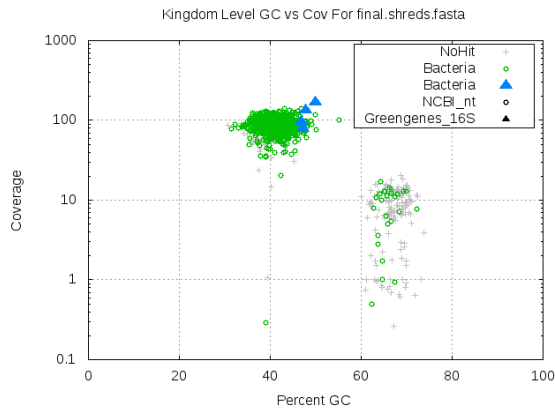


Species

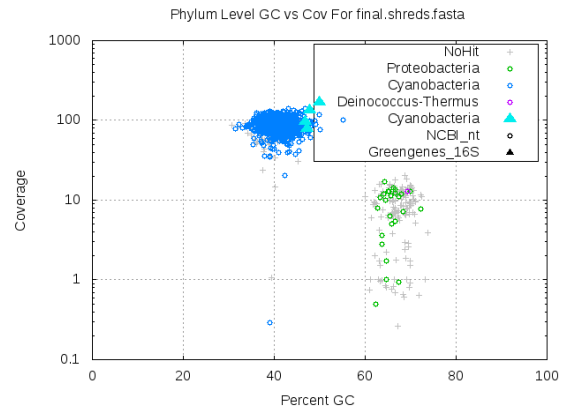


GC vs coverage based on GC of NCBI nt and Greengenes 16S rRNA gene hits to the assembly using megablast, shown for different taxonomic levels.

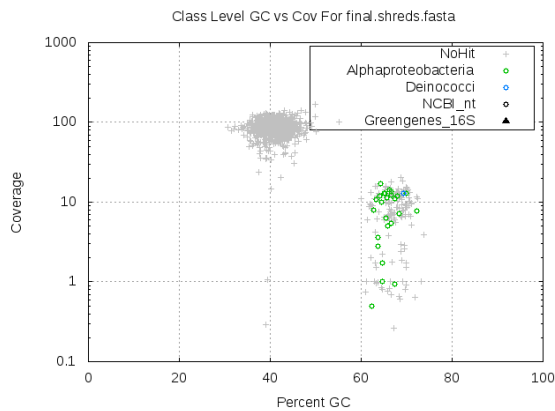
Kingdom



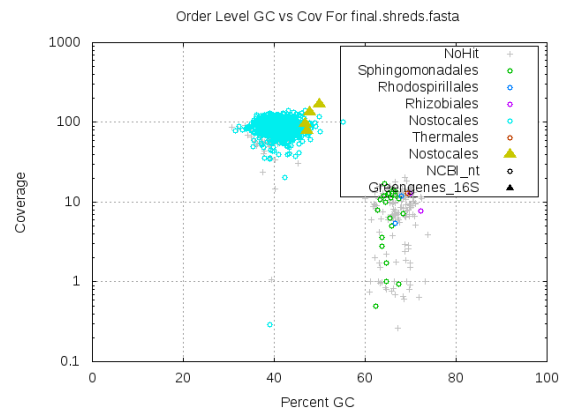
Phylum

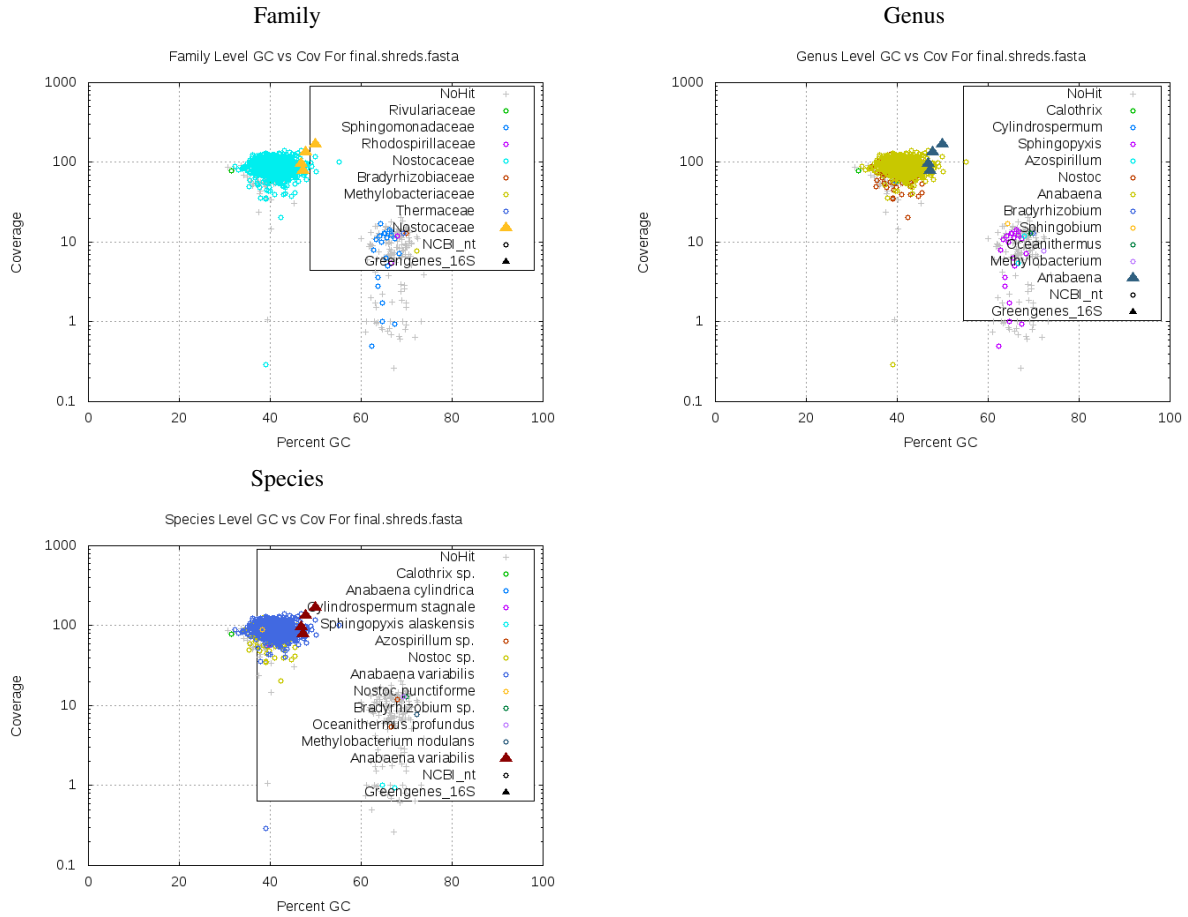


Class

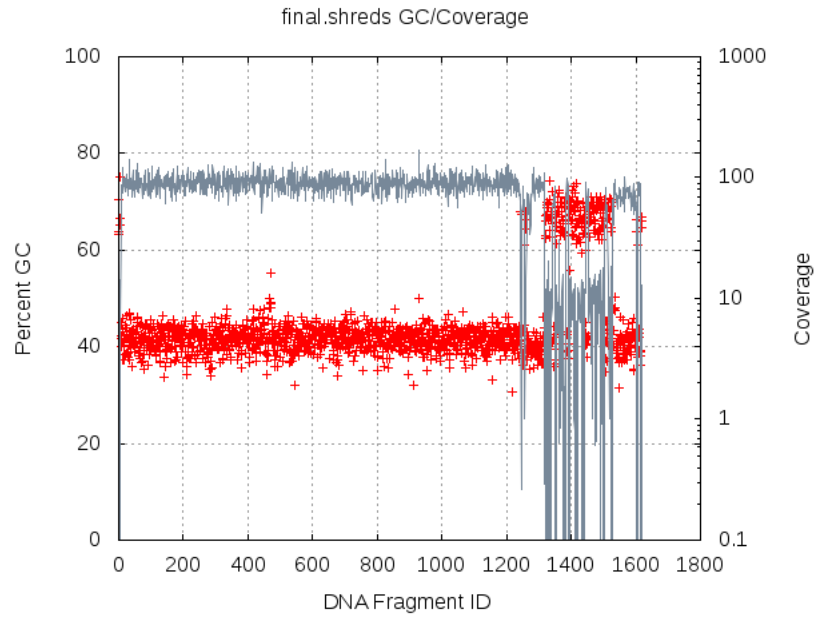


Order

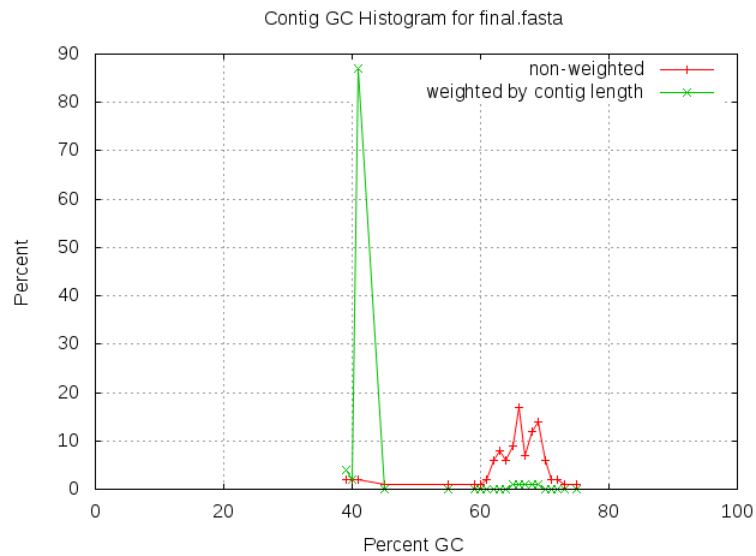




Coverage vs GC. Contigs were shredded into non-overlapping 5kbp and the GC of each shred was plotted as a point, colored by scaffold id. Coverage was calculated by mapping the fragment library to the final assembly and plotted as connected points.



GC histogram of the contigs, including contig length weighted distribution.



List of contigs and average percent GC bin:

Pct GC Bin	Contig Name
35	unitig_2 quiver, unitig_3 quiver, unitig_6 quiver
40	unitig_0 quiver, unitig_1 quiver, unitig_4 quiver, unitig_5 quiver, unitig_7 quiver, unitig_8 quiver, unitig_9 quiver
45	unitig_287 quiver
55	unitig_94 quiver, unitig_631 quiver
60	unitig_22 quiver, unitig_23 quiver, unitig_27 quiver, unitig_33 quiver, unitig_38 quiver, unitig_55 quiver, unitig_59 quiver, unitig_60 quiver, unitig_61 quiver, unitig_63 quiver, unitig_65 quiver, unitig_67 quiver, unitig_69 quiver, unitig_74 quiver, unitig_84 quiver, unitig_85 quiver,

	unitig_86 quiver, unitig_92 quiver, unitig_105 quiver, unitig_118 quiver, unitig_142 quiver, unitig_166 quiver, unitig_202 quiver, unitig_211 quiver, unitig_222 quiver, unitig_236 quiver, unitig_281 quiver, unitig_283 quiver, unitig_400 quiver, unitig_445 quiver, unitig_481 quiver, unitig_512 quiver, unitig_557 quiver, unitig_585 quiver, unitig_647 quiver, unitig_732 quiver, unitig_767 quiver, unitig_771 quiver, unitig_834 quiver, unitig_869 quiver
65	unitig_10 quiver, unitig_12 quiver, unitig_13 quiver, unitig_14 quiver, unitig_15 quiver, unitig_17 quiver, unitig_18 quiver, unitig_19 quiver, unitig_20 quiver, unitig_21 quiver, unitig_25 quiver, unitig_26 quiver, unitig_28 quiver, unitig_29 quiver, unitig_30 quiver, unitig_31 quiver, unitig_32 quiver, unitig_34 quiver, unitig_35 quiver, unitig_37 quiver, unitig_39 quiver, unitig_40 quiver, unitig_41 quiver, unitig_42 quiver, unitig_43 quiver, unitig_44 quiver, unitig_45 quiver, unitig_46 quiver, unitig_47 quiver, unitig_48 quiver, unitig_50 quiver, unitig_51 quiver, unitig_53 quiver, unitig_54 quiver, unitig_56 quiver, unitig_58 quiver, unitig_62 quiver, unitig_66 quiver, unitig_68 quiver, unitig_70 quiver, unitig_71 quiver, unitig_72 quiver, unitig_73 quiver, unitig_75 quiver, unitig_78 quiver, unitig_79 quiver, unitig_81 quiver, unitig_82 quiver, unitig_83 quiver, unitig_87 quiver, unitig_88 quiver, unitig_89 quiver, unitig_90 quiver, unitig_91 quiver, unitig_93 quiver, unitig_96 quiver, unitig_97 quiver, unitig_103 quiver, unitig_136 quiver, unitig_144 quiver, unitig_153 quiver, unitig_167 quiver, unitig_177 quiver, unitig_178 quiver, unitig_183 quiver, unitig_184 quiver, unitig_188 quiver, unitig_191 quiver, unitig_199 quiver, unitig_270 quiver, unitig_288 quiver, unitig_293 quiver, unitig_317 quiver, unitig_329 quiver, unitig_373 quiver, unitig_388 quiver, unitig_437 quiver, unitig_438 quiver, unitig_450 quiver, unitig_464 quiver, unitig_470 quiver, unitig_480 quiver, unitig_496 quiver, unitig_508 quiver, unitig_541 quiver, unitig_550 quiver, unitig_606 quiver, unitig_649 quiver, unitig_656 quiver, unitig_676 quiver, unitig_690 quiver, unitig_726 quiver, unitig_751 quiver, unitig_769 quiver, unitig_783 quiver, unitig_793 quiver, unitig_815 quiver, unitig_817 quiver, unitig_821 quiver, unitig_836 quiver, unitig_841 quiver, unitig_858 quiver, unitig_871 quiver, unitig_877 quiver, unitig_886 quiver
70	unitig_11 quiver, unitig_16 quiver, unitig_24 quiver, unitig_36 quiver, unitig_52 quiver, unitig_57 quiver, unitig_77 quiver, unitig_95 quiver, unitig_98 quiver, unitig_122 quiver, unitig_168 quiver, unitig_186 quiver, unitig_327 quiver, unitig_330 quiver, unitig_341 quiver, unitig_401 quiver, unitig_498 quiver, unitig_699 quiver, unitig_891 quiver
75	unitig_76 quiver

List of the top contig megablast hits against 16S ribosomal RNA genes.

Organism	Align Length (bp)	Pct Id	Contig Name
138867 <i>Anabaena variabilis</i> str. ATCC 29413 CP000117.1	1,491	100.00	unitig_0 quiver

5. Data Access

The following sequence fasta files can be downloaded from our JGI portal website.

<http://www.jgi.doe.gov/genome-projects>

The annotation of the assembled contigs can be found within IMG.

<http://img.jgi.doe.gov>

6. Methods

Isolate Improved Draft

Genome sequencing and assembly

The draft genome of *Nostoc sp.* was generated at the DOE Joint Genome Institute (JGI) using the Pacific Biosciences (PacBio) sequencing technology [1]. A >10kbp Pacbio SMRTbell™ library was constructed and sequenced on the PacBio RS platform, which generated 330,030 filtered subreads totaling 1.2 Gbp. All general aspects of library construction and sequencing performed at the JGI can be found at <http://www.jgi.doe.gov>. The raw reads were assembled using HGAP (version: 2.3.0, protocol version=2.3.0 method=RS.HGAP_Assembly.3, smrtpipe.py v1.87.139483,) [2]. The final draft assembly contained 178 contigs in 178 scaffolds, totaling 7.628 Mbp in size. The input read coverage was 136.9X.

Genome annotation

Genes were identified with Prodigal [3], followed by one round of manual curation using GenePRIMP [4] for genomes in fewer than 10 scaffolds. The predicted CDSs were translated and used to search the National Center for Biotechnology Information (NCBI) nonredundant database, UniProt, TIGRFam, Pfam, KEGG, COG, and InterPro databases.

The tRNAscanSE tool [5] was used to find tRNA genes, whereas ribosomal RNA genes were found by searches against models of the ribosomal RNA genes built from SILVA [6]. Other non-coding RNAs such as the RNA components of the protein secretion complex and the RNase P were identified by searching the genome for the corresponding Rfam profiles using INFERNAL [7]. Additional gene prediction analysis and manual functional annotation was performed within the Integrated Microbial Genomes (IMG) platform [8] developed by the Joint Genome Institute, Walnut Creek, CA, USA [9].

1. Eid John, et al. Real-Time DNA Sequencing from Single Polymerase Molecules. *Science* 2008
 2. Chin C, et al. Nonhybrid, finished microbial genome assemblies from long-read SMRT sequencing data. *Nat Methods* 2013
 3. Hyatt D, Chen GL, Lacascio PF, Land ML, Larimer FW, Hauser LJ. Prodigal: prokaryotic gene recognition and translation initiation site identification. *BMC Bioinformatics* 2010; 11:119.
 4. Pati A, Ivanova NN, Mikhailova N, Ovchinnikova G, Hooper SD, Lykidis A, Kyrpides NC. GenePRIMP: a gene prediction improvement pipeline for prokaryotic genomes. *Nat Methods* 2010; 7:455–457.
 5. Lowe TM, Eddy SR. tRNAscan-SE: a program for improved detection of transfer RNA genes in genomic sequence. *Nucleic Acids Res* 1997; 25:955–964.
 6. Pruesse E, Quast C, Knittel, Fuchs B, Ludwig W, Peplies J, Glckner FO. SILVA: a comprehensive online resource for quality checked and aligned ribosomal RNA sequence data compatible with ARB. *Nuc Acids Res* 2007; 35: 2188–7196.
 7. INFERNAL. Inference of RNA alignments. <http://infernal.janelia.org>.
 8. The Integrated Microbial Genomes (IMG) platform. <http://img.jgi.doe.gov>.
 9. Markowitz VM, Mavromatis K, Ivanova NN, Chen IMA, Chu K, Kyrpides NC. IMG ER: a system for microbial genome annotation expert review and curation. *Bioinformatics* 2009; 25:2271–2278.
-