riboSeed: leveraging prokaryotic genomic architecture to assemble across ribosomal regions

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Supplementary Data

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riboSeed (reference, riboSelect_clusters, reads, iters, flanking_width)
   ref = reference;
   clusters = parse riboSelect_clusters;
   region = clusters + flanking\_width;
   \mathbf{for}\ \mathit{iin\ iters\ do}
       map reads to ref;
       for cluster in clusters do
          filter and extract reads region;
          subassemble;
          return pseudocontig;
       end
       assess subassembly;
       if success then
          make pseudogenome from pseudocontigs;
          ref = pseudogenome;
       end
   end
   run assembler with reads and pseudocontigs;
```

Figure S1: Pseudocode of riboSeed algorithm

Making the artificial test genome

The artificial genome used for testing was constructed using the makeToyGenome.sh script included in the GitHub repository under the scripts directory. Briefly, the 7 rDNA regions from the *E. coli Sakai* genome were extracted with 5kb flanking sequence upstream and downstream; these sequences were then concatenated end to end to form a single, ~100kb sequence containing the 7 rDNAs as well as their flanking context.

Effect of reference sequence identity on riboSeed performance: Methods

A range of substitutions were introduced into a artificial genome using the runDegenerate.sh script (included in the GitHub repository under the scripts directory), which facilitates the following procedure. An artificial test genome is constructed (see above), and reads simulated using pIRS (100bp, 300bp inserts, stdev 10, 30-fold coverage, built-in

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Table S1: Hits resulting from searching the SRA database for various sequencing technologies as of January, 2017

Search term	Hits	Percentage
illumina	2242225	94.27
pacbio	21131	0.89
ion	30560	1.28
roche	42445	1.78
oxford	12301	0.52
solid	29791	1.25
Total	2378453	100

error profile). Then, for each of a range of substitution frequencies, substitutions are introduced into the simulated genome, either just in the flanking regions or throughout. riboSeed is run on the reads using the mutated genome as the reference, and the results are evaluated with riboScore. This script was run 100 times, using a different random seed each time. As the algorithm used by python's pseudo random number generator may differ between operating systems, comparable but not identical results can be expected.

Archaeal Datasets

We assessed the effectiveness of riboSeed with assembling archaeal genomes. Most (~55%) archaeal genomes have only a single rDNA, and none has been observed to have more than four. As riboSeed requires a sequencing dataset and a reference genome, applicability was limited; of the 104 entries in rrnDB with multiple rDNAs, only 7 had multiple entries at the species level. Among those, only 2 had publicly available short read data. We used riboSeed to reassemble Methanosarcina barkeri Fusaro DSMZ804 (Ion Torrent PGM, 89bp single-end reads) and Methanobacterium formicicum st. BRM9 (Illumina HiSeq 2000, 100bp paired-end reads). Methanobacterium formicicum st. JCM10132 (DRR017790) and Methanosarcina barkeri Fusaro DSMZ804 (SRR2064286) were the only ones that were suitable for riboSeed, meaning that there was publicly available short read data and that there is a related genome at the species level which is complete.

M. formicicum st. JCM10132 was sequenced on an Ion Torrent PGM, generating 106.5Mbp of single-end data. M formicicum BRM9 (CP006933.1) was used as a reference. The resulting de fere novo assembly resulted in assembly of 1 of 2 rDNA gaps. This represents the first application of riboSeed to Ion Torrent data.

Methanosarcina barkeri Fusaro DSMZ804 was sequenced using an Illumina HiSeq2000 with 101bp paired-end reads, with an average fragment length of 400bp. We downsampled to use 5% of the 19.4Gbp dataset. Methanosarcina barkeri str. Wiesmoor was used as a reference. The resulting riboSeed assembly showed correct assembly of 3 of 3 rDNAs, while de novo assemble failed to resolve any.

Taken together, we show that given appropriate datasets, archaeal datasets can be processed in the same manner used for bacteria.

Table S2: Accessions for 25 E. coli genomes

 $GCA_000021125.1_ASM2112v1$ $GCA_000023665.1_ASM2366v1$ $GCA_000026545.1_ASM2654v1$ $GCA_000262125.1_ASM26212v1$ $GCA_000273425.1_Esch_coli_MG12655_V1$ $GCA_000299255.1_ASM29925v1$ GCA_000714595.1_ASM71459v1 $GCA_000967155.1_HUSEC2011CHR1$ GCA_000974405.1_ASM97440v1 $GCA_000974465.1_ASM97446v1$ $GCA_000974575.1_ASM97457v1$ $GCA_001020945.2_ASM102094v2$ $GCA_001566675.1_ASM156667v1$ $GCA_002012245.1_ASM201224v1$ $GCA_001750845.1_ASM175084v1$ $GCA_001886755.1_ASM188675v1$ $GCA_001901145.1_ASM190114v1$ $GCA_002012145.1_ASM201214v1$ GCA_900096815.1_Ecoli_AG100_Sample2_M9_Assembly $GCA_002116715.1_ASM211671v1$ $GCA_002118095.1_ASM211809v1$ $GCA_002125925.1_ASM212592v1$ $GCA_001612475.1_ASM161247v1$ $GCA_001651965.1_ASM165196v1$ $GCA_001721125.1_ASM172112v1$

All available at ftp://ftp.ncbi.nlm.nih.gov/genomes/all/GCA/

Table S3: Strain names and accessions for reference genomes used in this study

Strain Name	Accession
E. coli MG1655 A. hydrophila ATCC 7966 B. cereus ATCC 10987 B. cereus NC7401 B. fragilis 638R R. sphaeroides ATCC 17029	NC_000913.3 NC_008570.1 AE017194.1 NC_016771.1 FQ312004.1 NC_009049.1, NC_009050.1
S. aureus TCH1516 S. aureus MRSA252 V. cholerae El Tor str. N16961 X. axonopodis pv. Citrumelo P. aeruginosa BAMCPA07-48 P. aeruginosa ATCC 15692	NC_010079.1 BX571856.1 NC_002505.1, NC_002506.1 CP002914.1 CP015377.1 NZ_CP017149.1

Table S4: Software Versions

Tool	Version
Mauve	2015-02-13 build 0
BLAST+ Barrnap	2.2.28+ 0.7
BWA	0.7.12-r1039
samtools	1.3.1
MAFFT SPAdes	v7.215 v3.9.0
QUAST	4.1
bedtools	2.17.0
EMBOSS	6.6.0
pIRS	2.0.2

BLAST Results for BA000007.2 rDNA

(Filtered to exclude matches less than 90% of query length and hits with E-value >10e-6)

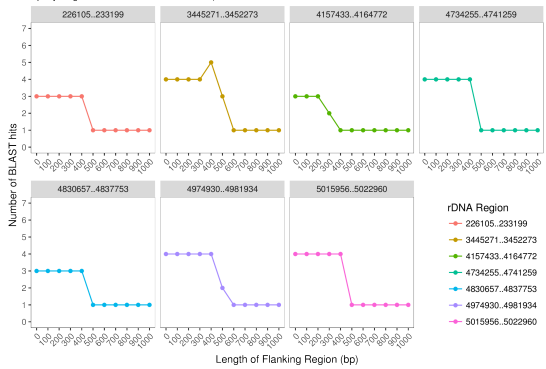


Figure S2: BLASTn was used to perform in silico DNA-DNA hybridization of all rDNA regions from $E.\ coli$ Sakai with variable flanking lengths. The number of hits is a proxy for occurrences in the genome; increasing the flanking length increases the specificity.

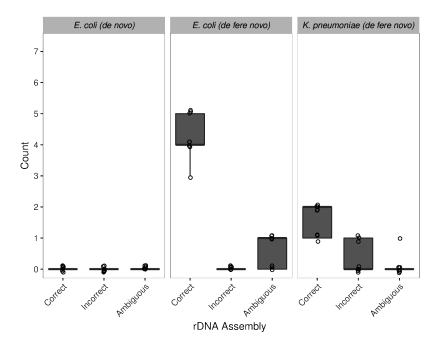
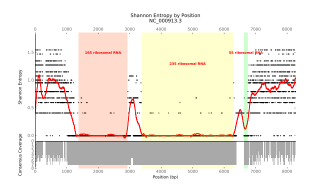
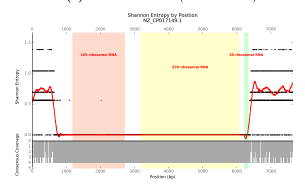


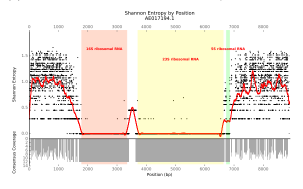
Figure S3: Assembly of artificial genome. De fere novo results in closure of 3-5 rDNAs with the correct reference; only 1-2 rDNAs are correctly assembled using K. pneumoniae. No rDNAs are assembled with de novo assembly. Scored with riboScore.py. N=8.



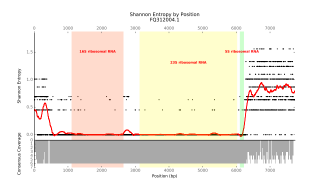
(a) E. coli MG1655 (NC_000913.3)



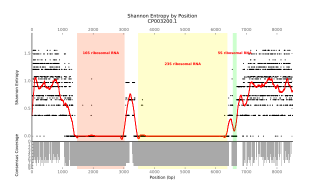
(c) P. aeruginosa strain ATCC 15692 (NZ_CP017149.1)



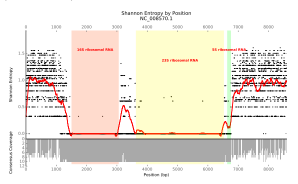
(e) B. cereus ATCC 10987 (AE017194.1)



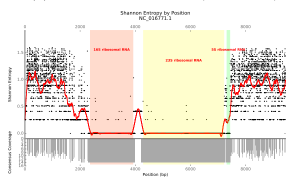
(g) B. fragilis 638R (FQ312004.1)



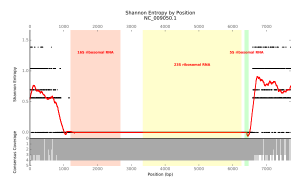
(b) K. pneumoniae subsp. pneumoniae HS11286 (CP003200.1)



(d) A. hydrophila ATCC 7966 (NC_008570.1)



(f) B. cereus NC7401 (NC_016771.1)



(h) R. sphaeroides ATCC 17029 (NC_009049.1, NC_009050.1)

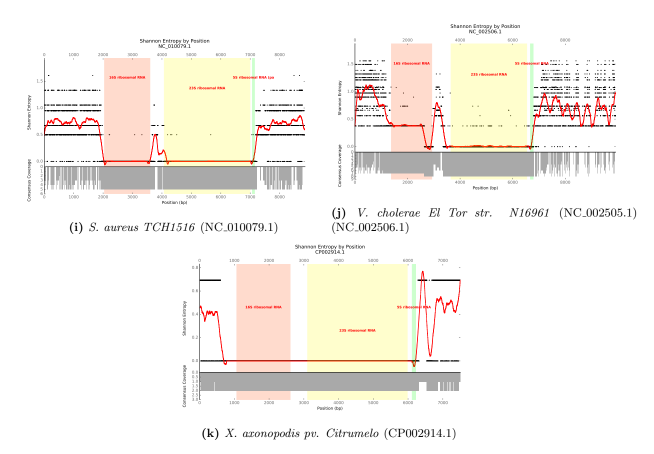


Figure S4: riboScan.py,riboSelect.py, and riboSnag.py were run on all the genomes used as references for *de fere novo* assemblies. Consensus alignment depth (grey bars) and Shannon entropy (black points, smoothed entropy as red line) for aligned rDNA regions.