DATA NOTE

The old and grumpy biogas metagenome

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Abstract

Background: a presentation of the interest or relevance of these data for the broader community

Findings: a very brief preview of the data type(s) produced, the methods used, and information relevant to data validation

Conclusions: a short summary of the potential uses of these data and implications for the field

Keywords: Biogas; Metagenome; Sequencing; Assembly; Annotation

Data description

Background

Biogas important energy source. Clean and awesome. Number of biogas plants in Germany/Europe/worldwide. Key is process optimization, not building more and more plants. Little is known about the microbial community responsible for everything.

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Here, we report the first deeply sequenced metagenome (and metatranscriptome?) of an agricultural production-scale biogas plant [1]. These data will enable a deeper exploration of the biogas-producting microbial community, a key step towards process optimization.

Digester management

The biogas plant, located in North Rhine Westphalia, Germany, features a mesophilic continuous wet fermentation technology and was designed for a capacity of $537\,kW_{el}$ combined heat and power (CHP) generation. The process comprises two digesters: a primary digester, where biogas and methane is produced, and a secondary digester, insignificant for production. The primary digester is fed hourly with a mixture of maize silage and liquid pig manure. The biogas and methane yields at the time of sampling were at 810.5 and 417.8 liters per kg organic dry matter $(l/kg\,oDM)$, respectively. After a theoretical retention time of 55 days, the digestate is stored in the closed, non-heated final storage tank. Further metadata are summarized in Table 1.

Sampling and DNA isolation

Samples from the primary digester of the aforementioned biogas plant were taken in November 2010. Prior to sampling process, approximately 15 L of fermenter substrate were discarded before aliquots of 1 L were transferred into clean gastight sampling vessels and transported directly to the laboratory.

A $20\,g$ aliquot of the fermentation sample was used for total community DNA preparation as described previously [2].

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Metagenomic sequencing

In total, we sequenced four different metagenome shotgun libraries with different insert sizes, resulting in over 23 gigabases scattered across 144 million reads. On Illumina's Genome Analyzer IIx system, we sequenced two libraries with an average insert size of $250\,nt$ and $450\,nt$, respectively, applying the Paired-End DNA Sample Preparation Kit (Illumina Inc.) as described by the manufacturer. On Illumina's MiSeq system, we sequenced two further libraries with an average insert size of $190\,nt$ and $690\,nt$, respectively, applying the Nextera DNA Sample Preparation Kit (Illumina Inc.) as described by the manufacturer. In all cases, we sequenced for approximately 300 cycles, applying the $2\times150\,nt$ paired-end protocol.

Eher kein 2x250: MiSeq v2 öffentlich 10/2012, Alex und ich hatten die MiSeq Daten in 07/2012!

Sequence quality control

Quality control with FastQC, version 0.11.2. Adapter contamination in Lane 7 (?). Adapter removal and quality trimming with Trimmomatic [3], version 0.32, using some paramters.

- 1 Remove adapters, using the included Truseq2-PE and Nextera-PE adapter files.
- 2 Remove leading and trailing low quality or N bases (below quality 3).
- 3 Adaptive quality trimming, balancing read length against error rate (target read length of 100 bases, strictness of 0.2, thus favouring read length over error sensitivity).
- 4 Drop reads below 36 bases long.

Either another table or brief summary post-QC.

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Needed?

k = 31

Metagenome assembly and quality assessment

2.3.1

We assembled the metagenome with Ray Meta [4], version 2.3.0, using a k-mer size of 31 and a minimum contig length of $1,000 \, bp$. This resulted in a total assembly size of approximately 217 megabases in 55,563 contigs, with an N50 value of $8,137 \, bp$. Table 2 summarizes our results.

2.2.4

LAP/ALE

We aligned the post-QC sequencing reads to the assembled contigs with bowtie2 [5], version 2.2.1, and used samtools [6], version 1.1, to convert SAM to BAM and thereafter sort the alignment file.

Gene prediction and annotation

We then used MetaProdigal [7], version 2.6.1, to predict 239,412 protein-coding genes on the assembled contigs. Table 3 also includes these results.

We blasted all predicted genes against the KEGG database [8], release 72.0, using Protein-Protein BLAST [9], version 2.2.29+. Of the 239,412 predicted genes, 230,354 were found in the KEGG database. We determined the KEGG Orthology (KO) for each gene by mapping the top-scoring BLAST hit to its KO.

Coverage analysis

0.6.1p2

We counted aligned reads in predicted genes with HTSeq [10], version 0.6.1p1. Figure 1 relates the metagenome and the metatranscriptome.

work in progress and up for discussion

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Availability

Data accession

The datasets supporting the results of this article are available in the [repository name] repository, [unique persistent identifier and hyperlink to datasets in http://format].

Reproducibility

The complete workflow is organized in a single GNU Makefile and available on GitHub [11]. Starting from the raw read files, available from SRA and/or GigaDB, all data and results can be reproduced by a simple invocation of *make*. Excluding KEGG analysis, which relies on a commercial license of the KEGG database, all steps are performed using free and open-source software.

phrasing

Requirements

The assembly of the metagenome data sets with Ray Meta requires 70 GB of RAM. Using only 40 cores, it completed within 12 hours wall-clock time. Annotation with blast took, of course, forever.

log runtime and memory in final run!

Discussion

Potential use cases. Metagenomic (and metatranscriptomic?) profiling of the biogasproducing microbial community. Identification of metaproteomic data out there (cite Vera, in preparation).

Ultimate goal: process optimization by biological insights.

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Competing interests

The authors declare that they have no competing interests

Author's contributions

AB conveived and performed all bioinformatic analyses and wrote the paper. IM investigated all metadata and drafted part of the data description. AP, ASch and AScz extensively revised the paper. ASch and AScz jointly directed the project. All authors read and approved the final manuscript.

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Figures

Figure 1 Relating the metagenome and metatranscriptome. Coverage or read counts, can discuss and decide once preliminary results are ready.

Tables

Table 1 Characteristics of the studied biogas plant. Primary digester, sampling date: Nov 15, 2010.

Process parameter	Sample
Net volume	$2041 m^3$
Dimensions	6.4m high, diameter of $21m$
Electrical capacity	$537 kW_{el}$
рН	7.83
Temperature	40 °C
Conductivity	22.10 mS/cm
Volative organic acids (VOA)	5327 mg/l
Total inorganic carbon (TIC)	14397mg/l
VOA/TIC	0.37
Ammoniacal nitrogen	2.93 g/l
Acetic acid	863mg/l
Propionic acid	76mg/l
Fed substrates	72% maize silage, $28%$ pig manure
Organic load	$4.0 kg oDM m^{-3} d^{-1}$
Biogas yield	810.5l/kgoDM
Methane yield	417.8 l/kg oDM

TIC only calculated, might be a few digits off

Table 2 Metagenome assembly. Some assembly statistics, minimum contig size of $1,000\,bp$.

Assembly metric	Our assembly
Total size	216,554,757 bp
Number of contigs	55,563
N50 value	8,137 bp
Largest contig	319,083 bp

Table 3 Gene prediction and annotation. Whatever.

Annotation metric	Our annotaion
Predicted genes	239,412
of these, full-length	160,124 (66.9%)
KEGG stuff	?

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Eher kein 2x250: MiSeq v2 öffentlich 10/2012, Alex und ich hatten die MiSeq	
Daten in 07/2012!	2
Needed? k=31	2
section not final!	2
2.3.1	2
2.2.4	2
LAP/ALE	2
$0.6.1 \mathrm{p2}$	2
work in progress and up for discussion	2
phrasing	3
log runtime and memory in final run!	3
section not final!	3
Funding Andreas S, Stadtwerke?	3
TIC only calculated, might be a few digits off	4