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allele.variability V.3

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Protocol status: Working

We use this protocol and it's working

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Abstract

A prerequisite to improve the predictability of microbial community dynamics is to understand their assembly mechanisms. To study factors that contribute to microbial community assembly, we examined temporal dynamics of genes in five aquatic metagenome time series, originating from marine off-shore or coastal sites and one lake, while focusing on a trait-based data evaluation.

We expected to find gene-specific patterns for the temporal allele variability depending on the metacommunity size of carrier-taxa and variability of the milieu and the substrates that the resulting enzymes are exposed to. In more detail we hypothesized that a larger metacommunity would cause increased temporal variability of functional units, as shown previously for taxonomic units. Furthermore, we hypothesized that multi-copy genes feature higher temporal variability than single-copy genes, because gene multiplication is often the consequence of increased variability in (subtil) changes of substrate quality and quantity. Finally, we hypothesized that direct exposure of proteins to the extracellular environment would result in increased temporal variability of the respective gene compared to intracellular proteins as they would be exposed to highly variable conditions. The first two hypotheses were confirmed in all, while an effect of the subcellular location of gene-products was only seen in three out of the five time series.

The gene with highest allele variability throughout all datasets was an iron transporter, which also represents a target for phage infections. This finding points to the general importance of iron transporter mediated phage infections on the assembly and maintenance of diversity of aquatic prokaryotes.

1 removal of nextera adaptors (cutadapt v1.8.3)

Command

Run for all raw data sequence revers and forward read-files from BATS and HOT

```
cutadapt -a CTGTCTCTTATA -o ca.file.forward.fastq.gz  
file.forward.fastq.gz  
cutadapt -a CTGTCTCTTATA -o ca.file.reverse.fastq.gz  
file.reverse.fastq.gz
```

2

Command

Run on BATS and HOT output files from step 1 and on LMO rawdata sequence files

```
sickle pe \  
-f /data/sara/LTG/pacific/BAT1/cutadapt/ca.file.forward.fastq.gz \  
-r /data/sara/LTG/pacific/BAT1/cutadapt/ca.file.reverse.fastq.gz \  
-t sanger \  
-o /data/sara/LTG/pacific/BAT1/sickle/qtrim.file.forward.fastq \  
-p /data/sara/LTG/pacific/BAT1/sickle/qtrim.file.reverse.fastq \  
-s /data/sara/gesifus.strains/sickle/qtrim.file.unpaired.fastq \  
-q 20 -l 50
```

3

Command

Run for SOLA (output files step 2). The personal reference database (personal_default_rRNA_DBs.fna) contains the silve rRNA sequences (downloaded 2013)

```
#interleave reads
merge-paired-reads.sh qtrim.file.forward.fastq
qtrim.file.reverse.fastq file.inter.fastq
#sortmerna
sortmerna --I file.inter.fastq --paired-in -n 2 --db
personal_default_rRNA_DBs.fna --other file.inter.protein -a 20
#unmerge protein data
unmerge-paired-reads.sh file.inter.protein.fastq
file.forward.protein.fastq file.reverse.protein.fastq
```

4

Command

Run for BATS, HOT (using output from step 2, only data from 10 sample days considered in this study) and SOLA (using output from step 3, data fro all available sample days)

```
fq2fa --merge --filter qtrim.file.1.fastq qtrim.file.2.fastq
file.merged.fa #interleave paired reads
cat file*.merged.fa > all.merged.fa #concatenate all individual
merged read files of a time series into one file
idba_ud --mink=25 --maxk=99 --step=4 -l all.merged.fa -o IDBAoutput
#run assembly
```

5

Command

Run for BATS, HOT and SOLA using the output file from step 4

```
prodigal -i IDBA.contig.fa -a prod.pep -d prod.fas -o prod.gff -f gff
```

6

Command

run for SOLA (predicted genes from step 5) and LMO (predicted genes from BARM assembly that were different from the public available data translated with table 11). The reference KEGG database (KEGG.faa) was downloaded on 15.05.2016.

```
makeblastdb -in KEGG.faa -parse_seqids -dbtype prot #create database  
for blastp  
blastp -db KEGG.faa -query prod.pep -outfmt 6 -num_alignments 1 -  
num_threads 16 -out blastout.tab #run blastp
```

7

Command

Diamond-Blast annotation (DIAMOND v0.8.22.84)

```
diamond makedb --in KEGG.faa -d KEGG #create diamond database  
diamond blastp -d KEGG -q prod.pep --more-sensitive -k 1 -o  
diamond.tab -p 26 #run diamond blast
```

8

Command

run for BATS, HOT, LMO (quality trimmed reads, step 3), SOLA (protein-coding reads, step 4) and MENDOTA (quality trimmed reads provided by collaborators). Assembled contigs were used as reference (BATS, HOT, SOLA: output from step 4; LMO: BARM assembly, MENDOTA: assembly provided by collaborators)

```
bowtie2-build contig.fa contigs #build indexed reference file  
bowtie2 --very-sensitive-local --no-unal -x contigs -1  
qtrim.file.forward.fastq -2 qtrim.file.reverse.fastq -S file.sam  
#for LMO, SOLA, BATS, HOT with paired-end reads  
bowtie2 --very-sensitive-local --no-unal -x contigs -U  
qtrim.file.fastq -S file.sam #for MENDOTA with single-end reads
```

9

Command

the prod.gff outputfile from step 5 was reformatted using a personal script (convert_prodigal_GFF_to_subread_featureCount_SAF.pl) to prod.saf concerning the requirements of the featureCounts software. For BATS, HOT, LMO, SOLA and MENDOTS sam-files produced during step 7 were used.

```
convert_prodigal_GFF_to_subread_featureCount_SAF.pl -g prod.gff -s  
prod.saf #converts prodigal output into prod.saf  
featureCounts -p -a prod.saf -T 30 -F SAF -o feature.tab *sam #for  
LMO, SOLA, BATS, HOT with paired-end reads  
featureCounts -a prod.saf -T 12 -F SAF -o feature.tab *sam #for  
MENDOTA with single-end reads
```

10

Command

Run for prokaryotic amino acid sequences from the KEGG database (downloaded Nov 2015) representing each KEGG ortholog (K*.prokaryotes.faa) with the alternative settings -n, -p or -a for sequences affiliating with gram negatives, gram positives or archaea, respectively

```
psort -n/p/a K*.prokaryotes.faa -o terse |awk -F'\t' '{$1="XXX" FS  
$1;}1' OFS='\t'|sed 1d > psort.K*.prok.tab #create psortoutput for  
each KEGG ortholog  
cat psort.K* >psort.all.prok.tab #concatenate results
```

11

Command

run for BATS, HOT, LMO, MENDOTA, SOLA: input files amino acid sequence files obtained after gene calling (prod.pep)

```
cd-hit -i prod.pep -o out.clust -c 1.00 -n 5 -M 16000 -T 22
```