# **Supplementary Material**

## A probabilistic model to recover individual genomes from metagenomes

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# **Supporting Data**

The simulated contigs, features files and scripts to reproduce the results are deposited under:

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# Supplementary Methods

## Poisson approximation for absolute abundance

When sequencing reads have been mapped to the contigs, we can quantify the number of reads that covers each position of each contig. This is the vector x with len(x) = L. We model the positional read coverage using a Poisson event model and assume that the positions are independent according to the Lander-Waterman statistics so that the joint likelihood is a product of positional likelihoods. Additionally, we scale the likelihood to a single event by taking the geometric mean. After simplification, the formula almost looks like the the Poisson over the mean contig coverage.

$$\mathcal{L}(\theta \mid \boldsymbol{x}) = \sqrt[L]{\prod_{i=1}^{L} \frac{\theta^{x_i}}{x_i!} e^{-\theta}} = \left(\frac{\prod_{i=1}^{L} \theta^{x_i}}{\prod_{i=1}^{L} x_i!} e^{-\theta L}\right)^{\frac{1}{L}} = \frac{\bar{\theta}}{\sqrt[L]{\prod_{i=1}^{L} x_i!}} e^{-\theta}$$
(1)

The data term in the denominator is a constant factor which is not dependent on  $\theta$ . It is the geometric mean over the  $x_i$ ! values which we approximate using the arithmetic mean  $\bar{x}$  of the positional contig coverage values.

$$\sqrt[L]{\prod_{i=1}^{L} x_i!} \approx \left(\frac{1}{L} \sum_{i=1}^{L} x_i\right)! = \bar{x}!$$
(2)

The approximation is good if the variance of the  $x_i$  is low. We use the approximation to avoid to handle other values than the mean which is usually computed. Since the term is a data constant, it is irrelevant for model comparison where only  $\theta$  differs among the genomes. The approximated likelihood using mean values is the standard Poisson formula.

$$\mathcal{L}'(\theta \mid \boldsymbol{x}) = \frac{\theta^{\bar{\boldsymbol{x}}}}{\bar{\boldsymbol{x}}!} e^{-\theta}$$
(3)

The log-likelihood is used in the MGLEX implementation for computational reasons. It is directly visible that the calculation is linear in the input.

$$\ell'(\theta \mid \boldsymbol{x}) = -\log \bar{\boldsymbol{x}}! + \bar{\boldsymbol{x}}\log\theta - \theta \tag{4}$$

#### **MLE for Poisson**

The multi-sample log-likelihood is the weighted sum over the sample log-likelihoods using mean vector  $a_i$  with length  $len(a_i) = M$ . This corresponds to the geometric mean in the exponential likelihood formula.

$$\ell(\boldsymbol{\theta} \mid \boldsymbol{a}_i) = \frac{1}{M} \sum_{j=1}^{M} -\log a_{i,j}! + a_{i,j} \cdot \log \theta_j - \theta_j$$
(5)

We select  $\boldsymbol{\theta}$  to maximize the joint log-likelihood  $f(\boldsymbol{\theta})$  on the training data a. The joint likelihood is a weighted sum of the log-likelihood values of all N contigs. Each contig's weight  $w_i$  is the contig length.

$$f(\boldsymbol{\theta}) = \sum_{i=1}^{N} w_i \cdot \ell(\boldsymbol{\theta} \mid \boldsymbol{a}_i) = \sum_{i=1}^{N} w_i \cdot \frac{1}{M} \sum_{j=1}^{M} -\log a_{i,j}! + a_{i,j} \log \theta_j - \theta_j$$
(6)

The partial derivative of f with respect to  $\theta_j$  for all  $j \in \{1...M\}$  is given by

$$\frac{\partial f}{\partial \theta_j} = \sum_{i=1}^N \frac{w_i}{M} \left( \frac{a_{i,j}}{\theta_j} - 1 \right) \tag{7}$$

We find the zeros of f to determine the MLE  $\hat{\theta}_j$ .

$$\sum_{i=1}^{N} \frac{w_i}{M} \left( \frac{a_{i,j}}{\theta_j} - 1 \right) = 0 \Leftrightarrow \sum_{i=1}^{N} \frac{w_i a_{i,j}}{\theta_j} = \sum_{i=1}^{N} w_i \Leftrightarrow \theta_j = \frac{\sum_{i=1}^{N} w_i a_{i,j}}{\sum_{i=1}^{N} w_i}$$
(8)

We see that the estimates for  $\theta_j$  maximize the joint log-likelihood because the second partial derivative with respect to  $\theta_j$  is always negative.

$$\frac{\partial^2 f}{\partial \theta_j^2} = -\sum_{i=1}^N \frac{w_i \, a_{i,j}}{M \, \theta_j^2} \tag{9}$$

### Binomial approximation for relative abundance

Similarly to the Poisson approximation for absolute abundance, we derive the Binomial approximation via a product of positional Binomials. Vector  $\boldsymbol{x}$  with length  $len(\boldsymbol{x}) = L$  holds the positional read coverage of a contig with length L for one sample and vector  $\boldsymbol{s}$  with same length holds the sum of positional read counts for the position i of the contig across all samples. There must be more than one sample to apply this model. We write the likelihood normalized to a single event as

$$\mathcal{L}(\theta \mid \boldsymbol{x}) = \sqrt[L]{\prod_{i=1}^{L} \binom{s_i}{x_i}} \theta^{x_i} (1-\theta)^{(s_i-x_i)}$$

$$= \sqrt[L]{\prod_{i=1}^{L} \binom{s_i}{x_i}} \cdot \sqrt[L]{\prod_{i=1}^{L} \theta^{x_i}} \cdot \sqrt[L]{\prod_{i=1}^{L} (1-\theta)^{(s_i-x_i)}}$$

$$= \sqrt[L]{\prod_{i=1}^{L} \binom{s_i}{x_i}} \cdot \theta^{\bar{x}} \cdot (1-\theta)^{(\bar{s}-\bar{x})}$$
(10)

The geometric mean of positional binomial coefficients (first term) is again a constant factor which is not dependent on  $\theta$ . We approximate this term using the arithmetic mean.

$$\sqrt[L]{\prod_{i=1}^{L} \binom{s_{i}}{x_{i}}} = \frac{\sqrt[L]{\prod_{i=1}^{L} s_{i}!}}{\sqrt[L]{\prod_{i=1}^{L} x_{i}! \cdot \sqrt[L]{\prod_{i=1}^{L} (s_{i} - x_{i})!}}}{\frac{1}{\sqrt[L]{\prod_{i=1}^{L} s_{i}!}} \approx \frac{\frac{1}{L} \sum_{i=1}^{L} s_{i}!}{\frac{1}{L} \sum_{i=1}^{L} x_{i}! \cdot \frac{1}{L} \sum_{i=1}^{L} (s_{i} - x_{i})!}} \qquad (11)$$

$$\approx \frac{\frac{1}{L} \sum_{i=1}^{L} s_{i}!}{\frac{1}{L} \sum_{i=1}^{L} s_{i}! \cdot \left(\frac{1}{L} \sum_{i=1}^{L} s_{i} - \frac{1}{L} \sum_{i=1}^{L} x_{i}\right)!} = \binom{\bar{s}}{\bar{x}}$$

The approximation is good if the differences in the coefficients are small. We use the approximation to avoid to handle other values than the mean which is usually computed. Since the term is a data constant, it is irrelevant for model comparison where only  $\theta$  differs among the genomes. The approximated likelihood using mean values is the standard Binomial formula.

$$\mathcal{L}'(\theta \mid \boldsymbol{x}) = \begin{pmatrix} \bar{s} \\ \bar{x} \end{pmatrix} \theta^{\bar{x}} (1-\theta)^{(\bar{s}-\bar{x})}$$
(12)

The log-likelihood is used in the MGLEX implementation for computational reasons. It is directly visible that the calculation is linear in the input.

$$\ell'(\theta \mid \boldsymbol{x}) = \log \begin{pmatrix} \bar{s} \\ \bar{x} \end{pmatrix} + \bar{x} \log \theta + (\bar{s} - \bar{x}) \log(1 - \theta)$$
(13)

#### **MLE for Binomial**

The multi-sample log-likelihood is the weighted sum over the sample log-likelihoods using mean vector  $\mathbf{r}_i$  with length  $len(\mathbf{r}_i) = M$ . This corresponds to the geometric mean in the exponential likelihood formula.

$$\ell(\boldsymbol{\theta} \mid \boldsymbol{r_i}) = \frac{1}{M} \sum_{j=1}^{M} \log \begin{pmatrix} R_i \\ r_{i,j} \end{pmatrix} + r_{i,j} \log \theta_j + (R_i - r_{i,j}) \log(1 - \theta_j)$$
(14)

 $R_i$  is the sum of the abundance vector  $r_i$ .

$$R_i = \sum_{j=1}^M r_{i,j} \tag{15}$$

Because both  $R_i$  and  $r_{i,j}$  can be real numbers, we need to generalize the binomial coefficient to positive real numbers via the gamma function  $\Gamma$ .

$$\log \binom{n}{k} = \log \Gamma(n+1) - \log \Gamma(k+1) - \log \Gamma(n-k+1)$$
(16)

We select  $\boldsymbol{\theta}$  to maximize the joint log-likelihood  $f(\boldsymbol{\theta})$  of the training data r. The joint likelihood is a weighted sum of the log-likelihood values of all N contigs. Each contig's weight  $w_i$  is the contig length.

$$f(\boldsymbol{\theta}) = \sum_{i=1}^{N} w_i \cdot \ell(\boldsymbol{\theta} \mid \boldsymbol{r_i})$$

$$= \sum_{i=1}^{N} w_i \cdot \frac{1}{M} \sum_{j=1}^{M} \log \binom{R_i}{r_{i,j}} + r_{i,j} \log \theta_j + (R_i - r_{i,j}) \log(1 - \theta_j)$$
(17)

The partial derivative of f with respect to  $\theta_j$  for all  $j \in \{1...M\}$  is given by

$$\frac{\partial f}{\partial \theta_j} = \sum_{i=1}^N \frac{w_i}{M} \left( \frac{r_{i,j}}{\theta_j} - \frac{R_i - r_{i,j}}{1 - \theta_j} \right) \tag{18}$$

We find the zeros of f to determine the MLE  $\hat{\theta}_{j}$ .

$$\sum_{i=1}^{N} \frac{w_i}{M} \left( \frac{r_{i,j}}{\theta_j} - \frac{R_i - r_{i,j}}{1 - \theta_j} \right) = 0$$

$$\Leftrightarrow (1 - \theta_j) \sum_{i=1}^{N} w_i r_{i,j} = \theta_j \left( \sum_{i=1}^{N} w_i R_i - \sum_{i=1}^{N} w_i r_{i,j} \right)$$

$$\Leftrightarrow \frac{1}{\theta_j} \sum_{i=1}^{N} w_i r_{i,j} = \sum_{i=1}^{N} w_i R_i$$

$$\Leftrightarrow \theta_j = \frac{\sum_{i=1}^{N} w_i r_{i,j}}{\sum_{i=1}^{N} w_i R_i}$$
(19)

Supplement v0.4.2p

We see that the estimates for  $\theta_j$  maximize the joint log-likelihood because the second partial derivative with respect to  $\theta_j$  is negative for our estimates  $\hat{\theta}_j$  for all  $j \in \{1...M\}$ .

$$\frac{\partial^2 f}{\partial \theta_j^2} = -\frac{R_i \theta_j^2 - 2r_{i,j} \theta_j + r_{i,j}}{(\theta_j - 1)^2 \theta_j^2}$$
(20)

$$-\frac{R_i\hat{\theta}_j^2 - 2r_{i,j}\hat{\theta}_j + r_{i,j}}{\left(\hat{\theta}_j - 1\right)^2\hat{\theta}_j^2} < 0 \Leftrightarrow \sum_{i=1}^N w_i r_{i,j} < \sum_{i=1}^N w_i R_i$$

$$\tag{21}$$

The last inequality is true by definition of  $R_i$  (assuming  $r_{i,j} \neq R_i$  for simplicity).

#### Naïve Bayes model for nucleotide composition

The Naïve Bayes model assumes independence of features so that the likelihood can be written as a product of likelihoods for all features. The feature vector  $\boldsymbol{x}$  for a contig contains nucleotide features such as all the absolute counts for all possible 5-mers. The length  $len(\boldsymbol{x})$  is M. The total sum of counts for the contig is S.

$$S = \sum_{i=1}^{M} x_i \tag{22}$$

The likelihood is normalized to a single event via the geometric mean.

$$\mathcal{L}(\boldsymbol{\theta} \mid \boldsymbol{x}) = \sqrt[S]{\prod_{i=1}^{M} \theta_i^{x_i}} = \prod_{i=1}^{M} \theta_i^{\frac{x_i}{S}} = \prod_{i=1}^{M} \theta_i^{x'_i}$$
(23)

Therefore, we directly use the normalized features.

$$x_i' = \frac{x_i}{\sum\limits_{j=1}^M x_j} \tag{24}$$

The log-likelihood is used in the MGLEX implementation for computational reasons. It is directly visible that the calculation is linear in the input.

$$\ell(\boldsymbol{\theta} \mid \boldsymbol{x'}) = \sum_{i=1}^{M} x'_i \log \theta_i$$
(25)

#### **MLE for Naive Bayes**

We select  $\boldsymbol{\theta}$  to maximize the joint log-likelihood  $f(\boldsymbol{\theta})$  on the training data c. The joint likelihood is a weighted sum of the log-likelihood values of all N contigs. Each contig's weight  $w_i$  is the contig length.

$$f(\boldsymbol{\theta}) = \sum_{i=1}^{N} w_i \cdot \ell(\boldsymbol{\theta} \mid \boldsymbol{c}_i) = \sum_{i=1}^{N} w_i \cdot \sum_{j=1}^{M} c_{i,j} \log \theta_j$$
(26)

Supplement v0.4.2p

We consider the constraint that  $sum(\theta) = 1$  because these are relative frequencies in each genome.

$$\sum_{j=1}^{M} \theta_j = 1 \tag{27}$$

Using the Lagrange method, we set up a function to maximize the joint data log-likelihood  $f(\theta)$  under the given constraint.

$$\Lambda(\boldsymbol{\theta}, \lambda) = f(\boldsymbol{\theta}) + \lambda \left( \left( \sum_{j=1}^{M} \theta_j \right) - 1 \right)$$
(28)

The partial derivative of  $\Lambda$  with respect to  $\theta_j$  for all  $j \in \{1...M\}$  is given by

$$\frac{\partial \Lambda}{\partial \theta_j} = \sum_{i=1}^N \frac{w_i c_{i,j}}{\theta_j} + \lambda \tag{29}$$

We find the zeros of  $\Lambda$  to determine the MLE  $\hat{\theta}_j$ .

$$\frac{\partial \Lambda}{\partial \theta_j} = 0 \Leftrightarrow \theta_j = \frac{\sum_{i=1}^N w_i c_{i,j}}{-\lambda}$$
(30)

Substituting  $\theta_j$  in Suppl. Equation 27 gives

$$-\lambda = \sum_{i=1}^{N} w_i \sum_{j=1}^{M} c_{i,j} = \sum_{i=1}^{N} w_i$$
(31)

The last simplification works because we work with normalized features that sum to one. Finally, we substitute  $-\lambda$  in (1) for the MLE.

$$\hat{\theta}_{j} = \frac{\sum_{i=1}^{N} w_{i} c_{i,j}}{\sum_{i=1}^{N} w_{i}}$$
(32)

#### Hierarchic Naive Bayes model for sequence similarity

We adapted the Naive Bayes model to weighted taxa by transforming the associated weights (i.e. alignments scores) into a set of sparse vectors  $x_l$ , one for each taxonomic rank. There are L such layers. The model likelihood is a product of observation probabilities, like in the standard Naive Bayes model, but the layers are also connected by multiplication.

$$\mathcal{L}(\boldsymbol{\theta} \mid \boldsymbol{x}) = \prod_{l=1}^{L} \prod_{j=1}^{len(\boldsymbol{x}_l)} \theta_{l,j}^{\boldsymbol{x}_{l,j}}$$
(33)

The small difference to the Naive Bayes model in the previous section is that there are no sequence length weights and that the feature vectors are not normalized. The multiplication of layers is a simplification because we know that taxonomic ranks are not independent. However, the model proved to be simple and effective for our purposes.

#### MLE for multi-layer Naive Bayes

Once the assumption of layer independence has been made, the problem simplifies to L independent Naive Bayes models with separate feature vectors and model parameters. The MLE derivation for each of these models is equivalent to the previous section.  $T_l$  is the number of features on level l.

$$\hat{\theta}_{l} = \frac{\sum_{i=1}^{N} t_{i,l}}{\sum_{j=1}^{T_{l}} \sum_{i=1}^{N} t_{i,l}}$$
(34)

## Metagenome simulation

We chose genomes according to the CAMI2015 (www.cami-challenge.org) medium complexity toy dataset which contained 450 different strains. Because some of the strains were simulated and had no accessible genome data, we reduced the dataset to 400 genomes with corresponding accessions. These comprised both finished and draft genomes. We sampled the abundance distributions from a lognormal with expectation value one and variance one, which produced abundance value in an reasonable range and formed relative abundance by normalization (Supplementary Table 1, column S1). We derived three secondary samples (Supplementary Table 1, columns S2, S3, S4) by separately applying continuous (exponential) growth to a randomly chosen set of genomes which each constituted 100 genomes (25%) in the primary sample using the following formula.

$$abundance'(genome) = abundance(gnome) \cdot 2^{growth\_rate(genome)}$$
 (35)

We modeled the change of the community composition in reaction to variation of environmental parameters, for instance if the growth medium is altered with no space restrictions then community members will grow according to their genomic potential. In our simplified growth model we choose the growth rate uniformly at random between one and ten regardless of the actual genome. We generated three secondary abundance profiles using the described procedure. We then simulated HiSeq Illumina reads for each sample using the ART simulator with read length 150 bp, insert size 270 bp and insert size standard deviation 27 bp. This corresponds to a common experimental setting because the reads are likely to overlap in the read assembly step. We chose a large yield of 15 Gb per sample to also cover genomes with low sample abundance (see Supplementary Table 1).

## Feature generation

All features are represented as separate text files, which can be compressed. Each line corresponds to a sequence but does not contain sequence identifiers. Therefore, it is required that the number and order of lines are identical in all features files.

## Sequence weights

We used the following GNU awk v4.0.1 script to calculate the length of each FASTA entry which we saved as contigs.seqlen.

```
#!/usr/bin/awk -f
BEGIN { id="\000" } # > not allowed in FASTA header
/^>/ {
    if( id != "\000" ) {
        printf "%s\t%s\n", id, sum;
    }
    id=substr( $0, 2 );
    sum = 0;
}
! /^>/ { sum+=length($0) }
END { printf "%s\t%s\n", id, sum }
```

## **5-mer frequencies**

We derived 5-mer frequencies for the gzip-compressed FASTA sequences using the program fasta2kmerS using the following GNU Bash syntax

```
zcat contigs.fna.gz |
fasta2kmersS -i <(cat) -f >(cat) -j 5 -k 5 -s 0 -h 0 -n 0 |
tr '\t' ' > contigs.kmc
```

#### **Taxonomic annotation**

We generated alignments using NCBI BLAST+/blastn v2.2.28+ in taxator-tk tabular format and filtered out all species level alignments using program alignments-filter from taxator-tk v1.3.3 which effectively removes the genomes of the same species from the reference sequences. Next we ran the program taxator with the LCA algorithm using only the best hits and processed the resulting GFF3 file. We used the alignment score as weight for each taxon and combined the annotations for each contig. Finally, we shortened the taxon paths using numbers and applied the described accumulation scheme to project alignment score onto higher-level taxa (see Table 1).

## Average read coverage

We aligned each sample's simulated read data to the artificial contigs with Bowtie v2.2.7 and converted the resulting SAM files to sorted BAM

```
bowtie2-build contigs.fna contigs.bowtie2
bowtie2 -x contigs.bowtie2 -1 forward.fq.gz -2 reverse.fq.gz |
samtools view -@ 5 -b - < input.sam | samtools sort -@ 5 - out</pre>
```

and then calculated the average read coverage using BedTools v2.25 and GNU awk v4.0.1

```
genomeCoverageBed -ibam out.sorted.bam -g contigs.seqlen -d -split |
awk 'BEGIN{IFS=0FS=FS="\t"}
{if($1 == last){ s+=$3; c+=1;}
else{if(s){print last, s/c; s=$3}; c=1; last=$1}}
END{print last, s/c}' > out.twocol.cov
```

Contigs which recruited no reads are omitted by BedTools, therefore zero values must be added afterwards by comparison to the sequence length file. Finally, we merged the coverage columns in Bash using

## **Performance measures**

In order to evaluate the quality of the predictions and to pick the optimal  $\beta$  parameter for the posterior estimation, MGLEX implements two measures: a mean squared error (MSE) and the mean pairwise coclustering (MPC) probability. Both require as input a label probability matrix which defines to which genome (column) each sequence (row) belongs, in terms of probabilities. In our simulation, the genome column corresponding to the source genome contained a one, all other columns a zero. A prediction probability matrix of the same form is required for comparison. In the case of ML predictions, this matrix also contains only ones and zeros and continuous values for the posterior estimation. Because sequences typically have different lengths, the user must provide a file with the sequence lengths (see AWK script for sequence weight file generation).

#### Mean squared error (MSE)

The mean squared error is the square root of the average squared difference between the label and the prediction matrix per contig (a value between zero and one). It is weighted by the length of the sequence.

$$MSE = \sqrt{\frac{1}{4\sum_{i=1}^{N} w_i} \sum_{i=1}^{N} w_i \sum_{j=1}^{M} (L_{i,j} - P_{i,j})^2}$$
(36)

Here, N is the number of sequences, M the number of genomes, w is a vector with the sequence lengths, L the label probability matrix and P the prediction probability matrix.

#### Mean pairwise coclustering (MPC)

The mean pairwise coclustering probability reports how likely a pair of sequences chosen from any genome among the real genomes, are found in the same predicted genome. The MPC averages over both, the pairs in the genomes and the genomes, regardless of their size. Since all sequences in our evaluations have the same length, we report the unweighted version of the MPC. The MPC is a probability between zero and one. It is easier to interpret than the MSE but requires more computation because it needs to consider all possible sequence pairs.

$$MPC = \frac{1}{|C|} \sum_{i=1}^{|C|} \left( \frac{1}{|C_i|(|C_i|-1)} \sum_{\substack{s_1, s_2 \in C_i \\ s_1 \neq s_2}} p(s_1|C_i) p(s_2|C_i) \right)$$
(37)

Here, the  $i^{\text{th}}$  genome is a set  $C_i$  which contains sequences  $s_i$  and C is a set which contains all genomes  $C_i$ .

#### Genome bin posterior

We calculate the bin posterior of a contig over the genome bins by normalization of the different likelihood values for each of the considered bins, so that their values sum to one. We assume, that the bin posterior is uniform over all G genome bins, so there is no additional weighting, for instance by genome size.  $\mathcal{L}(\text{genome} \mid \text{contig})$  is a vector which holds the likelihood of a specific contig for every genome bin. Then, the posterior is given by

$$P(\text{genome} \mid \text{contig}) = \frac{\mathcal{L}(\text{genome} \mid \text{contig})}{\sum\limits_{n=1}^{G} \mathcal{L}(\text{genome}_n \mid \text{contig})}$$
(38)

## Relative likelihood bin comparison

We derived a percentage similarity quantity S for two genome bins A and B, based on mixture likelihoods.

$$S(A,B) = \sqrt[z]{\prod_{i=1}^{N} \left(\frac{2L_i(\theta_A)L_i(\theta_B)}{L_i^2(\theta_A) + L_i^2(\theta_B)}\right)^{\frac{L_i^2(\theta_A) + L_i^2(\theta_B)}{L_i(\theta_A) + L_i(\theta_b)}}}$$
(39)

with normalization constant

$$Z = \sum_{i=1}^{N} \frac{L_{i}^{2}(\theta_{A}) + L_{i}^{2}(\theta_{B})}{L_{i}(\theta_{A}) + L_{i}(\theta_{b})}$$
(40)

Interestingly, when we interpret this quantity as a probability, a connection to the Kullback-Leibler divergence  $D_{\rm KL}$ , also called relative entropy, can be constructed. The Boltzmann formula (Suppl. Equation 41) establishes a general connection between entropy H and probability P.

$$\mathbf{H} = \log \mathbf{P} \tag{41}$$

When we substitute the probability P in Suppl. Equation 41 with S(A, B) from Suppl. Equation 39, we get

$$H(A, B) = -\frac{1}{Z} \sum_{i=1}^{N} \left( \frac{L_A^2 + L_B^2}{L_A + L_B} \right) \log \frac{L_i^2(\theta_A) + L_i^2(\theta_B)}{2 L_i(\theta_A) L_i(\theta_B)}$$

$$= -\frac{1}{Z} D_{\text{KL}}(\hat{L} \| L_{swap})$$
(42)

Suppl. Equation 42 is the negative Kullback-Leibler divergence over the sample data, which measures the loss of information when the suboptimal model with swapped parameters is used instead of the MLE parameter model, divided by the summed likelihood of the observed data.

# **Supplementary Tables**

Name	S1 (9)	so (%)	and 04.	SA (%)
		52 (%)	0.00	
Acaryochloris CCMEE 5410	0.27	0.07	0.08	80.0
Acetobacteraceae bacterium AI-5844	0.04	0.01	0.01	0.01
Acholeplasma laidlawii PG-8A	0.12	0.79	0.04	0.04
Acidaminococcus fermentans DSM 20731	0.16	0.04	0.05	0.05
Acidaminococcus BV3L6	0.29	0.08	0.21	0.92
Acidovorax ebreus TPSY	0.09	0.03	0.03	0.03
Acidovorax KKS102	0.21	0.96	1.23	0.06
Aciduliprofundum MAR08-339	1.12	0.31	0.34	0.34
Acinetobacter baumannii AB_TG2028	0.83	1.08	0.25	0.25
Acinetobacter baumannii Naval-113	0.13	0.25	0.18	0.04
Acinetobacter baumannii ZWS1122	0.05	0.06	0.01	0.01
Acinetobacter genomosp. 13TU NCTC 8102	0.06	0.02	0.02	0.12
Acinetobacter johnsonii ANC 3681	0.02	0.00	0.00	0.13
Acinetobacter nosocomialis 28F	0.07	0.02	0.02	0.02
Acinetobacter schindleri NIPH 900	0.01	0.00	0.05	0.06
Acinetobacter schindleri TG19614	0.08	0.20	0.34	0.32
Acinetobacter CIP 64.7	0.25	0.07	0.08	0.08
Actinobacillus minor NM305	0.23	0.06	0.07	0.07
Actinoplanes SE50/110	0.51	0.14	0.16	0.15
Actinopolyspora mortivallis DSM 44261	0.19	0.05	0.06	0.06
Aeromonas MDS8	0.16	0.04	0.05	0.29
Aggregatibacter actinomycetemcomitans AAS4A	0.02	0.00	0.01	0.01
Aggregatibacter actinomycetemcomitans SCC393	0.06	0.02	0.02	0.02
Alicyclobacillus acidocaldarius Tc-4-1	0.02	0.01	0.02	0.01
Alistipes CAG:53	0.14	0.04	0.28	0.04
Alloprevotella rava F0323	0.26	0.07	0.08	0.08
alpha proteobacterium LLX12A	0.07	0.02	0.02	0.02
alpha proteobacterium SCGC AAA015-019	0.04	0.15	0.01	0.35
alpha proteobacterium SCGC AAA536-G10	0.62	0.17	0.19	5.38
Alteromonas macleodii `Ionian Sea U8'	0.05	0.04	0.01	0.01
Amphibacillus xylanus NBRC 15112	0.10	0.03	0.09	0.03
Amycolatopsis mediterranei U32	0.07	0.02	0.02	0.02
Anaerococcus hydrogenalis ACS-025-V-Sch4	0.03	0.01	0.06	0.01
Anaerococcus hydrogenalis DSM 7454	0.18	0.05	0.06	0.06
Anaplasma marginale Florida	0.01	0.00	0.01	0.00
Anaplasma marginale Gypsy Plains	0.74	0.20	0.23	4.79
Anaplasma marginale St. Maries	0.52	0.14	4.64	0.16
Anoxybacillus SK3-4	0.15	0.04	0.05	0.05
Arthrobacter FB24	0.14	0.04	0.04	0.04
Arthrobacter TB 23	0.25	0.07	0.08	0.08
Azospirillum CAG:239	0.06	0.02	0.02	0.08
Bacillus amyloliquefaciens DC-12	0.34	0.09	0.11	0.10
Bacillus anthracis A0193	0.54	3.44	1.17	3.68
Bacillus anthracis A1055	0.16	0.10	0.05	0.05
Bacillus cereus Rock1-15	0.04	0.01	0.04	0.04
Bacillus cereus Rock4-2	0.30	0.23	0.09	0.09

Supplementary Table 1: Taxa in the simulated dataset and corresponding relative abundances for the primary sample S1 and the three secondary samples S2, S3 and S4.

Name	S1 (%)	S2 (%)	S3 (%)	S4 (%)
Bacillus cereus VD014	0.56	0.15	0.17	0.17
Bacillus pumilus ATCC 7061	0.14	0.37	0.04	0.04
Bacillus 37MA	0.14	0.04	0.20	0.04
Bacillus EGD-AK10	0.24	0.06	0.07	0.07
Bacillus WBUNB004	0.31	0.08	0.09	0.23
Bacillus WBUNB009	0.37	0.10	0.11	0.11
Bacillus subtilis gtP20b	0.14	0.04	0.98	0.15
Bacillus subtilis S1-4	0.46	0.13	0.14	0.14
Bacillus subtilis 6051-HGW	0.10	0.03	0.03	0.03
Bacillus thuringiensis BGSC 4CC1	0.12	0.03	0.04	0.04
Bacteriovorax DB6_IX	0.11	0.03	0.03	0.03
Bacteroides faecis CAG:32	0.06	0.05	0.33	0.02
Bacteroides fragilis CAG:558	0.08	0.06	0.03	0.03
Bacteroides 4_1_36	0.20	0.05	0.29	0.06
Bacteroides CAG:443	0.27	0.07	0.08	0.08
Bacteroides CAG:714	0.04	0.01	0.01	0.03
Beijerinckia indica ATCC 9039	0.06	0.02	0.22	0.07
Bifidobacterium longum CAG:69	0.02	0.02	0.01	0.01
Bizionia argentinensis JUB59	0.31	0.09	0.10	0.27
Bordetella bronchiseptica Bbr77	0.17	0.05	0.05	0.05
Borrelia burgdorferi 29805	0.48	0.13	0.15	0.15
Brachyspira hampsonii 30599	0.10	0.03	0.03	0.03
Bradyrhizobium DFCI-1	0.11	0.06	0.03	0.03
Bradyrhizobium S23321	0.30	2.08	0.09	0.09
Bradyrhizobium WSM2793	0.03	0.06	0.01	0.01
Brevibacillus laterosporus PE36	0.05	0.14	0.02	0.39
Brevibacterium casei S18	0.40	0.54	0.12	0.12
Brevibacterium mcbrellneri ATCC 49030	0.58	3.30	0.77	0.18
Brevundimonas abyssalis TAR-001	0.37	2.08	0.11	0.11
Brevundimonas BAL3	0.18	0.05	0.06	0.06
Brucella abortus 68-3396P	0.22	0.06	0.07	0.07
Brucella abortus NI274	0.17	0.25	0.20	0.05
Burkholderia bryophila 376MFSha3.1	0.04	0.01	0.01	0.01
Burkholderia mallei 2002721280	0.25	0.07	0.08	1.08
Burkholderia pseudomallei 668	0.16	1.10	0.05	0.05
Burkholderia pseudomallei DM98	0.13	0.04	0.04	0.04
Burkholderia CCGE1001	0.09	0.03	0.90	0.03
Burkholderia WSM4176	0.05	0.01	0.02	0.02
butyrate-producing bacterium SM4/1	0.27	2.17	0.08	0.08
Butyrivibrio crossotus CAG:259	0.07	0.02	0.06	0.02
Caldicellulosiruptor bescii DSM 6725	0.51	0.14	0.16	0.16
Caldivirga maquilingensis IC-167	0.06	0.02	0.02	0.02
Candidatus Accumulibacter phosphatis UW-1	0.25	0.07	0.08	0.08
Candidatus Photodesmus katoptron Akat1	0.20	0.57	0.06	0.22
Candidatus Poribacteria WGA-A3	0.06	0.02	0.02	0.02
Candidatus Saccharibacteria RAAC3_TM7_1	0.34	0.75	0.10	0.10
Capnocytophaga F0502	0.08	0.02	0.02	0.02
Carnobacterium WN1359	0.29	0.29	0.09	0.09

Name	S1 (%)	S2 (%)	S3 (%)	S4 (%)
Catellicoccus marimammalium M35/04/3	0.33	0.57	0.10	0.36
Chitinophaga pinensis DSM 2588	0.24	0.06	0.33	0.07
Chlamydia psittaci WC	0.05	0.01	0.02	0.20
Chlamydia trachomatis IU888	0.02	0.00	0.02	0.01
Chlamydia trachomatis L2b/Ams2	0.05	0.01	0.01	0.04
Chlamydia trachomatis RC-J/953	0.72	0.20	0.22	0.22
Chloroflexi bacterium oral isolate Chl1-2	0.35	0.09	0.11	0.11
Chloroflexi bacterium SCGC AB-629-P13	0.32	0.09	0.10	1.65
Citrobacter rodentium ICC168	0.08	0.02	0.02	0.02
Citrobacter KTE151	0.04	0.04	0.01	0.25
Clostridium acetobutylicum EA 2018	0.18	0.05	0.06	1.13
Clostridium carboxidivorans P7	0.23	0.25	2.18	0.07
Clostridium ATCC BAA-442	0.32	0.09	0.10	1.42
Clostridium CAG:269	0.38	0.10	0.83	1.36
Clostridium CAG:452	0.21	0.06	0.06	0.06
Clostridium CAG:567	0.44	0.12	0.14	0.53
Clostridium SY8519	0.10	0.03	0.03	0.03
Clostridium tyrobutyricum DSM 2637/ATCC 25755/JCM 11008	0.88	0.24	4.86	0.27
Collimonas fungivorans Ter331	0.19	0.05	0.33	0.06
Coprococcus comes CAG:19	0.10	0.03	0.09	0.03
Corynebacterium pseudotuberculosis 316	0.02	0.00	0.00	0.00
Corynebacterium pseudotuberculosis Cp162	0.08	0.02	0.02	0.20
Corynebacterium pseudotuberculosis I19	0.07	0.02	0.02	0.02
Corynebacterium KPL1855	0.82	4.06	0.25	0.60
Corynebacterium KPL1859	0.09	0.09	0.23	0.03
Corynebacterium KPL1998	0.09	0.03	0.03	0.26
Cronobacter sakazakii 701	0.11	0.03	0.03	0.03
Cupriavidus basilensis B-8	0.11	0.10	0.03	0.03
Cyanothece CCY0110	0.08	0.02	0.03	0.02
Cyclobacterium qasimii M12-11B	0.13	0.04	0.04	0.50
Desulfococcus oleovorans Hxd3	0.10	0.03	0.12	0.03
Desulfovibrio aespoeensis Aspo-2	0.22	0.06	0.07	0.07
Desulfurivibrio alkaliphilus AHT2	0.18	0.05	0.05	0.05
Dictyoglomus turgidum DSM 6724	0.39	0.11	0.12	0.12
Eggerthia catenaformis OT 569/DSM 20559	0.33	0.09	0.10	0.40
Emticicia oligotrophica DSM 17448	0.31	0.09	0.10	0.10
Enterobacter R4-368	0.07	0.02	0.02	0.02
Enterococcus flavescens ATCC 49996	0.08	0.02	0.02	0.02
Enterococcus GMD4E	1.14	0.31	0.35	0.35
Enterovibrio norvegicus FF-162	0.49	0.13	0.15	0.15
Erysipelotrichaceae bacterium 5_2_54FAA	0.27	0.15	0.08	0.08
Erythrobacter litoralis HTCC2594	0.86	0.23	0.26	0.26
Exiguobacterium pavilionensis RW-2	0.11	0.36	0.03	0.03
Facklamia ignava CCUG 37419	0.58	0.51	0.39	0.18
Faecalibacterium prausnitzii A2-165	0.08	0.02	0.02	0.02
Finegoldia magna BVS033A4	0.07	0.02	0.07	0.02
Firmicutes bacterium ASF500	0.09	0.02	0.03	0.03
Firmicutes bacterium CAG:170	0.17	0.05	0.05	0.05

Name	S1 (%)	S2 (%)	S3 (%)	S4 (%)
Fischerella thermalis PCC 7521	0.13	0.04	0.04	0.14
Flavobacteriaceae bacterium S85	0.41	0.11	0.13	0.13
Flavobacterium B17	0.14	0.04	0.04	0.04
Formosa AK20	0.64	0.18	0.20	0.20
Francisella tularensis 80700075	0.08	0.07	0.03	0.12
Frankia alni ACN14a	0.33	0.09	0.10	0.73
gamma proteobacterium IMCC2047	0.09	0.02	0.03	0.03
Gardnerella vaginalis 0288E	0.04	0.01	0.01	0.01
Gardnerella vaginalis 1500E	0.27	0.07	0.08	0.08
Geobacillus JF8	0.11	0.72	0.04	0.03
Gillisia marina	0.41	0.11	0.13	0.13
Glaciecola polaris LMG 21857	0.26	0.07	0.08	0.08
Glaciecola 4H-3-7+YE-5	0.33	0.21	0.10	0.10
Gordonia effusa NBRC 100432	0.09	0.03	0.45	0.03
Gordonia sihwensis NBRC 108236	0.12	0.24	0.04	0.11
Haemophilus aegyptius ATCC 11116	0.48	0.13	0.15	0.15
Haemophilus somnus 129PT	0.31	0.09	0.10	0.10
Haemophilus sputorum HK 2154	0.94	0.26	0.29	0.29
Haloferax BAB2207	0.43	0.12	0.13	0.13
Halomonas KM-1	0.13	0.03	0.04	1.02
Halorhabdus utahensis DSM 12940	0.01	0.00	0.04	0.00
Haloterrigena limicola JCM 13563	0.04	0.01	0.01	0.01
Helicobacter hepaticus ATCC 51449	0.38	0.10	3.30	0.39
Herbaspirillum B39	0.41	0.11	0.12	0.12
Ignavibacterium album JCM 16511	0.39	0.11	0.12	0.12
Isoptericola variabilis 225	0.20	0.05	0.06	0.06
Janibacter HTCC2649	0.43	0.12	0.13	0.95
Kingella kingae PYKK081	0.19	0.05	0.12	0.06
Klebsiella pneumoniae UHKPC01	0.18	1.40	0.06	0.06
Klebsiella pneumoniae UHKPCO2	0.14	0.04	1.04	0.04
Klebsiella pneumoniae UHKPC40	0.19	0.05	0.06	1.46
Ktedonobacter racemifer DSM 44963	0.14	0.04	0.04	0.04
Laceyella sacchari 1-1	0.08	0.02	0.02	0.21
Lachnospiraceae bacterium 2_1_46FAA	0.54	1.60	0.17	0.16
Lachnospiraceae bacterium 3-2	0.33	0.09	0.79	0.63
Lachnospiraceae bacterium 5_1_57FAA	0.04	0.22	0.01	0.37
Lachnospiraceae oral taxon 107 str. F0167	0.19	0.35	0.06	0.06
Lactobacillus acidipiscis KCTC 13900	0.04	0.01	0.01	0.01
Lactobacillus acidophilus 30SC	0.38	0.10	0.77	0.11
Lactobacillus acidophilus ATCC 4796	0.04	0.01	0.01	0.01
Lactobacillus casei 21/1	0.22	0.43	0.07	0.07
Lactobacillus casei Lpc-37	0.21	0.06	0.06	0.06
Lactobacillus delbrueckii ATCC BAA-365	0.03	0.01	0.01	0.01
Lactobacillus delbrueckii DSM 20072	0.32	0.09	1.90	2.76
Lactobacillus fermentum CECT 5716	0.42	0.11	0.13	0.13
Lactobacillus helveticus CNRZ32	0.08	0.20	0.02	0.02
Lactobacillus helveticus R0052	0.10	0.03	0.53	0.03
Lactobacillus iners ATCC 55195	0.06	0.02	0.02	0.06

Name	S1 (%)	S2 (%)	S3 (%)	S4 (%)
Lactobacillus iners LactinV 01V1-a	0.11	0.03	0.03	0.03
Lactobacillus plantarum 2165	0.85	0.23	0.26	0.54
Lactobacillus reuteri CF48-3A	0.66	0.18	0.20	0.20
Lactobacillus reuteri MM4-1A	0.23	1.14	0.07	0.07
Lactobacillus salivarius GJ-24	0.42	0.12	0.13	1.37
Lactobacillus ASF360	0.28	0.08	1.93	0.09
Legionella pneumophila str. 121004	0.05	0.06	0.01	0.29
Leifsonia xyli subxyli str. CTCB07	0.04	0.01	0.01	0.01
Leptospira borgpetersenii 200801910	0.20	0.05	0.06	0.06
Leptospira borgpetersenii 200901122	0.24	0.17	0.38	0.16
Leptospira interrogans Fiocruz R154	0.15	0.04	0.05	0.05
Leptospira interrogans L1207	0.11	0.03	0.04	0.03
Leptospira santarosai Oregon	0.59	3.58	0.18	0.18
Leptospira santarosai 2000027870	0.12	0.03	0.24	0.04
Leptospira santarosai HAI1380	0.13	0.04	0.04	0.04
Leuconostoc argentinum KCTC 3773	0.13	0.04	0.09	0.53
Leuconostoc citreum LBAE C10	0.02	0.01	0.20	0.07
Loktanella cinnabarina LL-001	0.23	0.06	0.07	0.07
Loktanella hongkongensis DSM 17492	0.18	0.05	0.06	0.53
Mannheimia haemolytica USDA-ARS-USMARC-183	0.19	0.05	0.06	0.06
marine gamma proteobacterium HTCC2080	0.10	0.06	0.30	0.03
Marinimicrobia bacterium SCGC AAA298-D23	0.26	0.65	0.08	0.08
Marinimicrobia bacterium SCGC AB-629-J13	0.26	0.07	0.08	0.08
Marinobacter EVN1	0.05	0.01	0.43	0.15
Megasphaera genomosp. type_1 str. 28L	0.14	0.04	0.47	0.04
Melissococcus plutonius DAT561	0.39	0.58	0.12	0.12
Mesoflavibacter zeaxanthinifaciens S86	0.11	0.03	0.21	0.03
Mesorhizobium LNHC229A00	0.16	1.18	0.11	0.05
Mesorhizobium LSHC416B00	0.04	0.01	0.01	0.01
Mesorhizobium LSJC264A00	0.04	0.08	0.01	0.01
Methanobrevibacter smithii TS146D	0.14	0.04	0.80	0.04
Methanobrevibacter smithii TS147C	0.14	0.04	0.04	0.11
Methanobrevibacter smithii TS95A	0.09	0.02	0.17	0.03
Methanocella arvoryzae MRE50	0.05	0.13	0.02	0.13
Methanosphaera stadtmanae DSM 3091	0.18	0.05	0.06	0.05
Methylobacterium extorquens PA1	0.10	0.78	0.03	0.03
Methyloglobulus morosus KoM1	0.19	0.05	0.06	1.18
Methylotenera versatilis 301	0.06	0.02	0.02	0.05
Methyloversatilis universalis EHg5	0.17	0.05	0.05	0.05
Microbacterium barkeri 2011-R4	0.12	0.03	0.04	0.04
Microbacterium 11MF	0.13	0.08	0.37	0.04
Microbacterium TS-1	0.10	0.03	0.03	0.03
Mobiluncus curtisii ATCC 43063	0.39	0.11	0.12	0.12
Mycobacterium abscessus 3A-0930-R	0.03	0.01	0.01	0.01
Mycobacterium abscessus 5S-0422	0.58	0.16	0.37	0.99
Mycobacterium abscessus M139	0.26	0.07	0.08	0.08
Mycobacterium chubuense NBB4	0.18	0.05	0.06	0.05
Mycobacterium intracellulare MOTT-02	0.19	0.05	0.06	0.12

Name	S1 (%)	S2 (%)	S3 (%)	S4 (%)
Mycoplasma gallisepticum NC08_2008.031-4-3P	0.05	0.02	0.02	0.02
Mycoplasma gallisepticum NY01_2001.047-5-1P	0.27	0.07	0.08	0.08
Neisseria gonorrhoeae PID18	0.13	0.03	0.04	0.04
Neisseria gonorrhoeae SK-92-679	0.13	0.04	0.04	0.04
Neisseria meningitidis NM1476	0.18	0.16	0.05	0.05
Neisseria meningitidis NM3223	0.16	0.05	0.05	0.15
Neisseria meningitidis NM604	0.27	0.07	0.08	0.08
Neisseria sicca 4320	0.09	0.23	0.03	0.03
Niabella aurantiaca DSM 17617	0.15	0.23	0.32	0.04
Nitrolancea hollandica Lb	0.36	0.78	1.24	0.11
Nocardia tenerifensis NBRC 101015	0.40	0.11	0.12	1.67
Nocardiopsis CNS639	0.74	0.20	0.23	0.23
Nonomuraea coxensis DSM 45129	0.69	1.26	0.21	1.07
Oceanicaulis HTCC2633	0.19	0.05	0.34	0.06
Oceanobacillus kimchii X50	0.74	2.17	0.23	0.86
Octadecabacter arcticus 238	0.06	0.02	0.02	0.02
Paenibacillus alvei TS-15	0.19	0.05	1.36	0.06
Paenibacillus larvae BRL-230010	0.03	0.01	0.01	0.01
Paenibacillus Aloe-11	0.04	0.01	0.01	0.01
Pantoea AS-PWVM4	0.09	0.02	0.03	0.03
Parabacteroides ASF519	0.19	0.05	0.06	0.85
Parascardovia denticolens IPLA 20019	0.42	0.11	2.48	3.38
Parasutterella excrementihominis CAG:233	0.72	0.20	0.22	1.60
Patulibacter americanus DSM 16676	0.07	0.02	0.02	0.55
Patulibacter medicamentivorans	0.45	0.12	1.09	0.14
Pediococcus acidilactici D3	0.07	0.05	0.02	0.02
Pelosinus fermentans A11	0.04	0.01	0.23	0.03
Peptoclostridium difficile P20	0.12	0.03	0.04	0.53
Peptoclostridium difficile P48	0.04	0.01	0.01	0.08
Peptoclostridium difficile P53	0.24	0.07	1.05	0.07
Polynucleobacter necessarius QLW-P1DMWA-1	0.09	0.02	0.03	0.03
Porphyromonas gingivalis JCVI SC001	0.17	0.05	0.23	1.25
Porphyromonas gingivalis W50	0.81	0.22	0.25	0.25
Porphyromonas macacae DSM 20710/JCM 13914	0.11	0.03	0.03	0.03
Prevotella salivae DSM 15606	0.04	0.01	0.01	0.01
Prevotella C561	0.03	0.19	0.01	0.01
Prevotella CAG:1185	0.39	0.11	3.30	0.12
Prevotella CAG:592	0.35	1.04	1.14	0.11
Prevotella CAG:617	0.32	0.09	0.10	0.91
Prevotella CAG:755	0.14	0.04	0.04	0.04
Prevotella CAG:873	0.07	0.02	0.02	0.02
Pseudomonas aeruginosa BWHPSA006	0.10	0.03	0.03	0.03
Pseudomonas aeruginosa LESB58	0.23	0.20	1.02	0.07
Pseudomonas aeruginosa PABL056	0.09	0.03	0.03	0.03
Pseudomonas mendocina ymp	0.10	0.03	0.03	0.18
Pseudomonas CF161	0.07	0.02	0.02	0.02
Pseudomonas EGD-AK9	0.04	0.01	0.03	0.01
Pseudomonas M47T1	0.28	0.08	0.09	0.39

Pseudomonas TJI-51         0.03         0.01         0.01         0.01           Pseudomonas syringae pv. lachrymans M302278         0.25         0.70         1.23         0.08           Psychobactur PRuf-1         0.03         0.01         0.01         0.01           Pyrobaculum aerophilum str. IN2         0.32         0.09         0.10         0.07           Pyrobaculum calidifontis JCM 11548         0.03         0.01         0.01         0.07           Pyrococcus furiosus COMI         0.36         0.10         0.11         0.11           Raltochering salmoniarum ATCC 33209         0.49         0.13         0.15         0.15           Rhizobium BRG74         0.07         0.12         0.02         0.01         0.01         0.01           Rhodobacter SW2         0.17         0.05         0.05         0.05         0.05         0.05         0.05         0.06         0.05         0.05         Not         0.04         0.13         0.14         2.85           Rhodobacter SW2         0.17         0.70         0.22         0.02         0.02         0.02         0.02         0.02         0.02         0.02         0.02         0.02         0.02         0.02         0.02         0.02	Name	S1 (%)	S2 (%)	S3 (%)	S4 (%)
Pseudononas syringas pv. lachrymans M302278         0.25         0.70         1.23         0.08           Psychrobacter PRwf-1         0.03         0.01         0.01         0.01           Pyrobaculum calidifontis JCM 11548         0.03         0.01         0.01         0.01           Pyrobaculum calidifontis JCM 11548         0.03         0.01         0.01         0.01           Pyrobaculum calidifonts JCM 11548         0.03         0.01         0.01         0.01           Ralstonia solanacearum Po82         0.18         0.05         0.06         0.05           Renibacterium salmoninarum ATCC 33209         0.49         0.13         0.15         0.15           Rhizobium IRBG74         0.07         0.12         0.02         0.70           Rhodobacter sphaeroides ATCC 17029         0.47         0.13         0.14         2.85           Rhodococcus Subdrilla MS 552         0.22         0.66         0.07         0.07           Rhodococcus Subdrilla Sultics SWK14         0.55         0.17         0.17           Rhodospeudomonas palustris BisB5         0.28         0.08         0.08           Rhodospeudomonas palustris Sultis         0.22         0.02         0.02           Rhodospeudomonas palustris BisB5         0.28 <td>Pseudomonas TJI-51</td> <td>0.03</td> <td>0.01</td> <td>0.01</td> <td>0.01</td>	Pseudomonas TJI-51	0.03	0.01	0.01	0.01
Psychocbacter PMer-1       0.03       0.01       0.01       0.01         Pyrobaculum calidifontis JCM 11548       0.32       0.09       0.10       0.10         Pyrobaculum calidifontis JCM 11548       0.33       0.01       0.01       0.01         Pyrocaccus furiosus CDM1       0.36       0.10       0.11       0.11         Ralstonia solanacarum P682       0.18       0.02       0.01       0.01       0.01         Rhizobium phaseoli (h74-10)       0.08       0.33       0.02       0.02       0.02         Rhizobium Phaseoli (h74-10)       0.08       0.33       0.02       0.07         Rhodobacter sphaeroides ATCC 17029       0.47       0.13       0.14       2.85         Rhodobacter aphaeroides ATCC 17029       0.47       0.12       0.02       0.02         Rhodobacter aphaeroides ATCC 17029       0.42       1.34       0.07       0.22         Rhodobacter aphaeroides ATCC 17029       0.42       1.34       0.07       0.22         Rhodobacter aphaeroides ATCC 17029       0.42       1.34       0.07       0.22         Rhodobacter aphaeroides ATCC 17029       0.42       1.3       0.17       0.12         Rhodobacter aphaeroides ATCC 17029       0.42       1.10	Pseudomonas syringae pv. lachrymans M302278	0.25	0.70	1.23	0.08
Pyrobaculum aerophilum str. IM2       0.32       0.09       0.00       0.01       0.01         Pyrobaculum calidifontis JCM 11548       0.03       0.01       0.01       0.01         Ralstonia solanacearum P682       0.18       0.05       0.06       0.05         Ranibacterium salmoninarum ATCC 33209       0.49       0.13       0.15       0.15         Rhizobium phaseoli Ch24-10       0.08       0.33       0.02       0.02         Rhodobacter SW2       0.17       0.05       0.05       0.05         Rhodobacter subactise ATCC 17029       0.47       0.13       3.28       0.42         Rhodobacter subactise ATCC 17029       0.47       0.13       3.28       0.42         Rhodobacter subactise SW11       1.39       3.41       3.28       0.42         Rhodobacter SW2       0.24       1.34       0.07       0.22         Rhodosoccus Pholni LMG 5362       0.24       1.34       0.07       0.22         Rhodosoccus Pholni LMG 5462       0.24       1.34       0.67       0.70         Rhodosoccus P27       0.22       0.66       0.67       0.06       0.68         Rhodosocus P27       0.22       0.68       0.40       0.70       0.17 <t< td=""><td>Psychrobacter PRwf-1</td><td>0.03</td><td>0.01</td><td>0.01</td><td>0.01</td></t<>	Psychrobacter PRwf-1	0.03	0.01	0.01	0.01
Pyrobaculum calidifontis JCM 11548         0.03         0.01         0.01         0.01           Pyrococcus furiosus COM1         0.36         0.10         0.11         0.11           Ralstonia solanacearum Po82         0.18         0.05         0.06         0.05           Renibacterium salmoninarum ATCC 33209         0.49         0.13         0.01         0.01           Rhizobium phaseoli Ch24-10         0.08         0.33         0.02         0.70           Rhodobacter sphaeroides ATCC 17029         0.17         0.05         0.05         Rhodobacter sphaeroides ATCC 17029         0.41         0.34         0.07         0.22           Rhodobacter sphaeroides ATCC 17029         0.47         0.15         0.17         0.22           Rhodobacter aphaeroides ATCC 17029         0.47         0.22         0.00         7         0.22           Rhodococcus 29MTsu3.1         0.06         0.02         0.02         0.02         0.02         0.02         0.02           Rhodopsirillum rubrum ATCC 1170         0.22         0.66         0.68         Rickettsia rickettsii str. 'Sheila Smith'         0.49         0.65         0.66         0.66           Rickettsia rickettsii str. 'Sheila Smith'         0.49         0.61         0.17         0.17	Pyrobaculum aerophilum str. IM2	0.32	0.09	0.10	0.10
Pyrcoccus furiosus GM1         0.36         0.10         0.11         0.11           Ralstonia solanacearum Po82         0.18         0.05         0.06           Renibacterium salmoninarum ATCC 33209         0.49         0.13         0.15         0.15           Rnizobium etli Brasil 5         0.02         0.01         0.01         0.01           Rnizobium IRBG74         0.08         0.33         0.02         0.02           Rhodobacter SW2         0.17         0.05         0.05         0.05           Rhodobacter SW2         0.47         0.13         0.14         2.85           Rhodobacter SW2         0.47         0.13         0.14         2.85           Rhodobacter SW2         0.24         1.34         0.42         0.02         0.02           Rhodococcus Phodnii LMC 5362         0.24         1.34         0.70         0.22           Rhodosoccus Phodnii LMC 5362         0.24         0.06         0.07         0.07           Rhodosoccus Phodnii LMC 5362         0.22         0.02         0.02         0.01         0.01           Rhodosoccus Phodnii LMC 5362         0.26         0.06         0.06         0.06           Rhodosoccus Phodnii LMC 51817         0.22         0.21	Pyrobaculum calidifontis JCM 11548	0.03	0.01	0.01	0.07
Ralstonia solanacearum Po82       0.18       0.05       0.06       0.05         Renibacterium salmoninarum ATCC 33209       0.49       0.13       0.15       0.15         Rhizobium phaseoli Ch24-10       0.08       0.33       0.02       0.02         Rhizobium IRBG74       0.07       0.12       0.02       0.70         Rhodobacter SW2       0.17       0.05       0.05       0.05         Rhodobacter sphaeroides ATCC 17029       0.47       0.13       0.14       2.85         Rhodobacter scae bacterium KLH1       1.39       3.41       3.28       0.42         Rhodococcus 29MFTsu3.1       0.06       0.02       0.02       0.07         Rhodopseudononas palustris BisE5       0.28       0.08       0.08       0.08         Rhodopsirillum rubrum ATCC 11170       0.02       0.01       0.01       0.05         Rickettsia helvetica C9P9       0.19       0.55       0.66       0.66         Rickettsia helvetica C9P9       0.90       0.32       1.97       1.27         Remerella anatipestifer RA-YM       0.08       0.02       0.03       0.65         Rudaella lutea DSN 19387       0.72       0.20       0.22       0.22       0.22       0.22       0.22	Pyrococcus furiosus COM1	0.36	0.10	0.11	0.11
Renibacterium salmoninarum ATCC 33209       0.49       0.13       0.15       0.15         Rhizobium etli Brasil 5       0.02       0.01       0.01         Rhizobium phaseoli Ch24-10       0.08       0.33       0.02       0.02         Rhizobium IRBG74       0.07       0.12       0.02       0.70         Rhodobacter sphaeroides ATCC 17029       0.47       0.13       0.14       2.85         Rhodobacter sphaeroides ATCC 17029       0.47       0.13       0.14       2.85         Rhodobacteraceae bacterium KLH11       1.39       3.41       3.28       0.42         Rhodocccus P20FTSU3.1       0.06       0.02       0.02       0.02         Rhodopirellula baltica SWK14       0.55       0.15       0.17       0.17         Rhodospeudononas palustris BisB5       0.28       0.08       0.08       Rhodes         Rickettsia helvetica CSP9       0.19       0.05       0.06       0.06         Rickettsia nicketsis istr. Sheila Smith'       0.42       0.11       0.13       0.13         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Ruminiclostridium thermocellum YS       0.55       0.15       1.98       0.17         R	Ralstonia solanacearum Po82	0.18	0.05	0.06	0.05
Rhizobium etli Brasil 5       0.02       0.01       0.01       0.01         Rhizobium phaseoli Ch24-10       0.08       0.33       0.02       0.02         Rhizobium INBG74       0.07       0.12       0.02       0.70         Rhodobacter SW2       0.17       0.05       0.05       0.05         Rhodobacter sphaeroides ATCC 17029       0.47       0.13       0.14       2.85         Rhodobacterscaee bacterium KLH1       1.39       3.41       3.28       0.42         Rhodocccus PMFTNU3.1       0.06       0.02       0.02       0.02         Rhodopseudomonas palustris BisB5       0.28       0.08       0.08       0.08         Rhodopseudomonas palustris BisB5       0.28       0.06       0.06       0.06         Rickettsi helvetica C9P9       0.19       0.05       0.06       0.06         Rickettsi natricketsi str. "Sheila Smith"       0.49       0.32       1.97       1.27         Riemerella anatipestifer RA-YM       0.08       0.02       0.03       0.65         Ruminiclostridium thermocellum YS       0.55       0.15       1.98       0.17         Ruminococcus CAG:352       0.10       0.03       0.03       0.03         Ruminococcus CAG:352	Renibacterium salmoninarum ATCC 33209	0.49	0.13	0.15	0.15
Rhizobium Phaseoli Ch24-10       0.08       0.33       0.02       0.02         Rhizobium IRBG74       0.07       0.12       0.02       0.70         Rhodobacter SW2       0.17       0.05       0.05       0.05         Rhodobacter Synaeroides ATCC 17029       0.47       0.13       0.14       2.85         Rhodobacteraceae bacterium KLH11       1.39       3.41       3.28       0.42         Rhodocccus PMFTsu3.1       0.06       0.02       0.02       0.02         Rhodopirellula baltica SWK14       0.55       0.15       0.17       0.17         Rhodopirellula baltica SWK14       0.55       0.16       0.06       0.06         Rickettsia rickettsii str.       Shella Smith'       0.49       0.32       1.97       1.27         Riemerella anatipestifer RA-YM       0.08       0.02       0.03       0.65         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13	Rhizobium etli Brasil 5	0.02	0.01	0.01	0.01
Rhizobium IRBG74         0.07         0.12         0.02         0.70           Rhodobacter SW2         0.17         0.05         0.05         0.05           Rhodobacter spharoides ATCC 17029         0.47         0.13         0.14         2.85           Rhodobacters caee bacterium KLH11         1.39         3.41         3.28         0.42           Rhodocccus Podnii LMC 5362         0.24         1.34         0.07         0.22           Rhodocccus P27         0.22         0.06         0.07         0.07           Rhodopseudomonas palustris BisB5         0.28         0.08         0.08         0.08           Rhodoseillum rubrum ATCC 11707         0.02         0.01         0.05         0.06         0.06           Rickettsia helvetica C9P9         0.19         0.05         0.06         0.06         0.02         0.22	Rhizobium phaseoli Ch24-10	0.08	0.33	0.02	0.02
Rhodobacter SW2       0.17       0.05       0.05       0.05         Rhodobacter sphaeroides ATCC 17029       0.47       0.13       0.14       2.85         Rhodobacter sphaeroides ATCC 17029       0.47       0.13       0.14       2.85         Rhodobacter sphaeroides ATCC 17029       0.24       1.34       0.07       0.22         Rhodocccus 29MTsu3.1       0.06       0.02       0.02       0.02         Rhodopseudomonas palustris BisB5       0.28       0.08       0.08       0.08         Rhodospirillum rubrum ATCC 11170       0.02       0.01       0.01       0.05         Rickettsia rickettsii str. Sheila Smith'       0.49       0.32       1.97       1.27         Riemerella antipestifer RA-YM       0.08       0.02       0.02       0.22         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Ruminicoccus CAG:382       0.10       0.03       0.03       0.03         Ruminococcus CAG:379       0.88       2.32       0.27       0.27         Salinispora arenicola CNY330       0.52       0.72       0.16       0.16         Salinonella enterica SA-2       0.05       0.01       0.02       0.02         S	Rhizobium IRBG74	0.07	0.12	0.02	0.70
Rhodobacter sphaeroides ATCC 17029       0.47       0.13       0.14       2.85         Rhodobacteraceae bacterium KLH11       1.39       3.41       3.28       0.42         Rhodcocccus 29WTFu3.1       0.06       0.02       0.02       0.02         Rhodcoccus 29WTFu3.1       0.06       0.02       0.02       0.02         Rhodcoccus P27       0.22       0.06       0.07       0.07         Rhodospirillum baltica SW14       0.55       0.15       0.17       0.17         Rhodospirillum rubrum ATCC 11170       0.02       0.01       0.01       0.05         Rickettsia helvetica C9P9       0.19       0.05       0.06       0.06         Ruickettsia rickettsii str. 'Sheila Smith'       0.49       0.32       1.97       1.27         Ruimiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Ruminiclostridium thermocellum YS       0.55       0.15       1.98       0.17         Ruminococcus CAG:822       0.10       0.03       0.03       0.03         Ruminococcus CAG:82       0.27       0.27       0.26       0.22         Salinispora arenicola CNY234       0.61       0.17       0.19       0.19         Salinispora parifi	Rhodobacter SW2	0.17	0.05	0.05	0.05
Rhodobacteraceae bacterium KLH11       1.39       3.41       3.28       0.42         Rhodococcus rhodnii LMG S362       0.24       1.34       0.07       0.22         Rhodococcus 29MFTsu3.1       0.06       0.02       0.02       0.02         Rhodopseudomonas palustris BisB5       0.28       0.08       0.08       0.08         Rhodospirillum rubrum ATCC 11170       0.19       0.05       0.06       0.07         Rickettsia helvetica CSP9       0.19       0.05       0.06       0.06         Rickettsia rickettsii str. 'Sheila Smith'       0.49       0.32       1.97       1.27         Riemerella anatipestifer RA-TM       0.08       0.02       0.01       0.13       0.15         Rudanella lutea DSM 19387       0.72       0.20       0.22       0.22         Ruminiclostridium thermocellum YS       0.55       0.15       1.98       0.17         Ruminococcus CAG:579       0.88       2.32       0.27       0.27         Salinispora arenicola CNY244       0.61       0.04       0.04       0.04         Salinispora arenicola CNY234       0.61       0.17       0.19       0.19         Salinispora arenicola CNY234       0.61       0.17       0.10       0.22	Rhodobacter sphaeroides ATCC 17029	0.47	0.13	0.14	2.85
Rhodococcus rhodnii LMG 5362       0.24       1.34       0.07       0.22         Rhodococcus 29MTru3.1       0.06       0.02       0.02       0.02         Rhodopcoccus 29T       0.22       0.06       0.07       0.07         Rhodoprellula baltica SWK14       0.55       0.15       0.17       0.17         Rhodopsirillum rubrum ATCC 11170       0.02       0.01       0.01       0.05         Rickettsia rickettsii str.       Sheila Smith'       0.49       0.32       1.97       1.27         Riemerella anatipestifer RA-YM       0.08       0.02       0.03       0.65         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Ruminiclostridium thermocellum YS       0.55       0.15       1.98       0.17         Ruminococcus CAG:382       0.10       0.03       0.03       0.03         Salinispora arenicola CNT849       0.86       0.23       0.26       0.26         Salinispora arenicola CNT849       0.61       0.17       0.19       0.19         Salinonella enterica SA-2       0.05       0.01       0.02       0.02         Salmonella enterica SA-2       0.05       0.01       0.02       0.02         <	Rhodobacteraceae bacterium KLH11	1.39	3.41	3.28	0.42
Rhodococcus 29MFTsu3.1       0.06       0.02       0.02       0.02         Rhodococcus P27       0.22       0.06       0.07       0.07         Rhodopirellula baltica SWK14       0.55       0.15       0.17       0.17         Rhodospirillum rubrum ATCC 11170       0.02       0.01       0.01       0.05         Rickettsia helvetica C9P9       0.19       0.05       0.06       0.06         Rickettsia rickettsii str. 'Sheila Smith'       0.49       0.32       1.97       1.27         Riemerella anatipestifer RA-YM       0.08       0.02       0.03       0.65         Rudanella lutea DSM 19387       0.72       0.20       0.22       0.22         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Rumincoccus CAG:362       0.10       0.03       0.03       0.03         Ruminococcus CAG:579       0.88       2.32       0.27       0.27         Salinispora arenicola CNY234       0.61       0.17       0.19       0.19         Salinispora arenicola CNY330       0.52       0.72       0.16       0.16         Salmonella enterica CFSAN001588       0.13       0.04       0.94       0.04         Selenomonas noxia AT	Rhodococcus rhodnii LMG 5362	0.24	1.34	0.07	0.22
Rhodococcus P27       0.22       0.06       0.07       0.07         Rhodopirellula baltica SWK14       0.55       0.15       0.17       0.17         Rhodopseudomonas palustris BisB5       0.28       0.08       0.08       0.08         Rhodospirillum rubrum ATCC 11170       0.02       0.01       0.01       0.05         Rickettsia helvetica C9P9       0.19       0.05       0.06       0.06         Rickettsia rickettsii str. `Sheila Smith'       0.49       0.32       1.97       1.27         Riemerella anatipestifer RA-YM       0.08       0.02       0.02       0.22       0.22         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13       0.13         Ruminiclostridium thermocellum YS       0.55       0.15       1.98       0.17         Ruminococcus CAG:32       0.10       0.03       0.03       0.03         Saccharomonospora cyanea NA-134       0.13       0.04       0.04       0.04         Salinispora arenicola CNT849       0.66       0.23       0.26       0.26         Salinispora arenicola CNT849       0.61       0.17       0.19       0.19         Salinispora arenicola CNT330       0.52       0.72       0.16	Rhodococcus 29MFTsu3.1	0.06	0.02	0.02	0.02
Rhodopirellula baltica SWK14       0.55       0.15       0.17       0.17         Rhodopseudomonas palustris BisB5       0.28       0.08       0.08         Rhodospirillum rubrum ATCC 11170       0.02       0.01       0.01       0.05         Rickettsia helvetica C9P9       0.19       0.05       0.06       0.06         Rickettsia rickettsii str. `Sheila Smith'       0.49       0.32       1.97       1.27         Riemerella anatipestifer RA-YM       0.08       0.02       0.03       0.65         Rudanella lutea DSM 19387       0.72       0.20       0.22       0.22         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Ruminococcus CAG:382       0.10       0.03       0.03       0.03         Ruminococcus CAG:579       0.88       2.32       0.27       0.27         Sacharomonospora cyanea NA-134       0.13       0.04       0.04       0.04         Salinispora arenicola CNY234       0.61       0.17       0.19       0.19         Salinispora pacifica CNY330       0.52       0.72       0.16       0.16         Salmonella enterica SA-2       0.05       0.01       0.02       0.02       0.22         Se	Rhodococcus P27	0.22	0.06	0.07	0.07
Rhodopseudomonas palustris BisB5       0.28       0.08       0.08       0.08         Rhodospirillum rubrum ATCC 11170       0.02       0.01       0.01       0.05         Rickettsia helvetica C9P9       0.19       0.05       0.06       0.06         Rickettsia rickettsii str. `Sheila Smith'       0.49       0.32       1.97       1.27         Riemerella anatipestifer RA-YM       0.08       0.02       0.02       0.22       0.22         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Ruminococcus CAG:382       0.10       0.03       0.03       0.03         Ruminococcus CAG:579       0.88       2.32       0.27       0.27         Saccharomonospora cyanea NA-134       0.13       0.04       0.04       0.04         Salinispora arenicola CNT849       0.86       0.23       0.26       0.26         Salinispora arenicola CNY234       0.61       0.17       0.19       0.19         Salmonella enterica CFSAN001588       0.13       0.04       0.04       0.04         Selenomonas noxia ATCC 43541       0.06       0.02       0.02       0.02         Shigella dysenteriae CDC 74-1112       0.20       1.73       0.55       0.58<	Rhodopirellula baltica SWK14	0.55	0.15	0.17	0.17
Rhodospirillum rubrum ATCC 11170       0.02       0.01       0.01       0.05         Rickettsia helvetica C9P9       0.19       0.05       0.06       0.06         Rickettsia rickettsii str. `Sheila Smith'       0.49       0.32       1.97       1.27         Riemerella anatipestifer RA-YM       0.08       0.02       0.02       0.22       0.22         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Ruminococcus CAG:382       0.10       0.03       0.03       0.03         Ruminococcus CAG:579       0.88       2.32       0.27       0.27         Sacharomonospora cyanea NA-134       0.13       0.04       0.04       0.04         Salinispora arenicola CNT349       0.86       0.23       0.26       0.26         Salinispora arenicola CNT330       0.52       0.72       0.02       0.02         Salmonella enterica SA-2       0.05       0.01       0.02       0.02         Salmonella enterica CFSAN001588       0.13       0.04       0.04       0.01         Shigella boydii 965-58       0.18       0.05       0.58       0.05         Shigella flexneri 1485-80       0.11       0.03       0.03       0.03 <tr< td=""><td>Rhodopseudomonas palustris BisB5</td><td>0.28</td><td>0.08</td><td>0.08</td><td>0.08</td></tr<>	Rhodopseudomonas palustris BisB5	0.28	0.08	0.08	0.08
Rickettsia helvetica C9P9       0.19       0.05       0.06       0.06         Rickettsia rickettsii str. 'Sheila Smith'       0.49       0.32       1.97       1.27         Riemerella anatipestifer RA-YM       0.08       0.02       0.03       0.65         Rudanella lutea DSM 19387       0.72       0.20       0.22       0.22         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Ruminiclostridium thermocellum YS       0.55       0.15       1.98       0.17         Ruminococcus CAG:382       0.10       0.03       0.03       0.03         Ruminococcus CAG:579       0.88       2.32       0.27       0.27         Salcharomonospora cyanea NA-134       0.13       0.04       0.04       0.04         Salinispora arenicola CNT349       0.86       0.23       0.26       0.26         Salinispora pacifica CNY330       0.52       0.72       0.16       0.16         Salmonella enterica SA-2       0.05       0.01       0.02       0.02         Salenomas noxia ATCC 43541       0.06       0.02       0.02       0.02         Shewanella frigidimarina NCIMB 400       0.04       0.04       0.04       0.04	Rhodospirillum rubrum ATCC 11170	0.02	0.01	0.01	0.05
Rickettsia rickettsii str. 'Sheila Smith'       0.49       0.32       1.97       1.27         Riemerella anatipestifer RA-YM       0.08       0.02       0.03       0.65         Rudanella lutea DSM 19387       0.72       0.20       0.22       0.22         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Rumincoccus CAG:382       0.10       0.03       0.03       0.03         Rumincoccus CAG:579       0.88       2.32       0.27       0.27         Saccharomonospora cyanea NA-134       0.13       0.04       0.04       0.04         Salinispora arenicola CNT849       0.86       0.23       0.26       0.26         Salinispora pacifica CNY330       0.52       0.72       0.16       0.16         Salmonella enterica SA-2       0.05       0.01       0.02       0.02         Salmonella enterica CFSAN001588       0.13       0.04       0.89       0.04         Selenomonas noxia ATCC 43541       0.06       0.02       0.02       0.02         Shigella bygiti 965-58       0.18       0.05       0.58       0.05         Shigella flexneri 1485-80       0.11       0.03       0.03       0.03         Simonsiella mu	Rickettsia helvetica C9P9	0.19	0.05	0.06	0.06
Riemerella anatipestifer RA-YM       0.08       0.02       0.03       0.65         Rudanella lutea DSM 19387       0.72       0.20       0.22       0.22         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Ruminiclostridium thermocellum YS       0.55       0.15       1.98       0.17         Ruminococcus CAG:382       0.10       0.03       0.03       0.03         Ruminococcus CAG:579       0.88       2.32       0.27       0.27         Saccharomonospora cyanea NA-134       0.13       0.04       0.04       0.04         Salinispora arenicola CNT849       0.86       0.23       0.26       0.26         Salinispora pacifica CNY330       0.52       0.72       0.16       0.16         Salmonella enterica SA-2       0.05       0.01       0.02       0.02         Salmonella enterica CFSAN001588       0.13       0.04       0.94       0.04         Selenomonas noxia ATCC 43541       0.06       0.02       0.02       0.02         Shigella flexneri 1485-80       0.11       0.03       0.03       0.03         Shigella flexneri 1485-80       0.11       0.03       0.03       0.03         Shigella flexneri	Rickettsia rickettsii str. `Sheila Smith'	0.49	0.32	1.97	1.27
Rudanella lutea DSM 19387       0.72       0.20       0.22       0.22         Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Ruminiclostridium thermocellum YS       0.55       0.15       1.98       0.17         Ruminococcus CAG:382       0.10       0.03       0.03       0.03         Ruminococcus CAG:579       0.88       2.32       0.27       0.27         Saccharomonospora cyanea NA-134       0.13       0.04       0.04       0.04         Salinispora arenicola CNT849       0.61       0.17       0.19       0.19         Salinispora pacifica CNY330       0.52       0.72       0.16       0.16         Salmonella enterica SA-2       0.05       0.01       0.02       0.02         Salenomonas noxia ATCC 43541       0.06       0.02       0.02       0.02         Shigella boydii 965-58       0.18       0.05       0.55       0.45         Shigella flexneri 2930-71       0.11       0.03       0.03       0.03         Shigella flexneri 2930-71       0.11       0.03       0.03       0.03         Shigella flexneri 2930-71       0.11       0.03       0.03       0.03         Shigella flexneri 1485-80	Riemerella anatipestifer RA-YM	0.08	0.02	0.03	0.65
Ruminiclostridium thermocellum ATCC 27405       0.42       0.11       0.13       0.13         Ruminiclostridium thermocellum YS       0.55       0.15       1.98       0.17         Ruminococcus CAG:382       0.10       0.03       0.03       0.03         Ruminococcus CAG:579       0.88       2.32       0.27       0.27         Saccharomonospora cyanea NA-134       0.13       0.04       0.04       0.04         Salinispora arenicola CNT849       0.66       0.23       0.26       0.26         Salinispora arenicola CNY330       0.52       0.72       0.16       0.16         Salmonella enterica SA-2       0.05       0.01       0.02       0.02         Salmonella enterica CFSAN001588       0.13       0.04       0.04       0.04         Shewanella frigidimarina NCIMB 400       0.04       0.04       0.01       0.13       0.03       0.03         Shigella bydii 965-58       0.18       0.05       0.58       0.05       0.11       0.03       0.03       0.03         Shigella flexneri 1485-80       0.11       0.03       0.03       0.03       0.03       0.03       0.03       0.03       0.03       0.03       0.03       0.03       0.03       0.03 <t< td=""><td>Rudanella lutea DSM 19387</td><td>0.72</td><td>0.20</td><td>0.22</td><td>0.22</td></t<>	Rudanella lutea DSM 19387	0.72	0.20	0.22	0.22
Ruminiclostridium thermocellum YS       0.55       0.15       1.98       0.17         Ruminococcus CAG:382       0.10       0.03       0.03       0.03         Ruminococcus CAG:579       0.88       2.32       0.27       0.27         Saccharomonospora cyanea NA-134       0.13       0.04       0.04       0.04         Salinispora arenicola CNT849       0.86       0.23       0.26       0.26         Salinispora arenicola CNY234       0.61       0.17       0.19       0.19         Salmonella enterica SA-2       0.05       0.01       0.02       0.02         Salmonella enterica CFSAN001588       0.13       0.04       0.04       0.04         Selenomonas noxia ATCC 43541       0.06       0.02       0.02       0.02         Shigella boydii 965-58       0.18       0.05       0.58       0.05         Shigella flexneri 1485-80       0.11       0.03       0.03       0.03         Shigella flexneri 2930-71       0.11       0.08       0.09       0.09         Sphingopyxis MC1       0.07       0.07       0.02       0.08         Streptococcus agalactiae GB00264       0.09       0.03       0.03       0.03         Streptococcus agalactiae MRI Z1-022	Ruminiclostridium thermocellum ATCC 27405	0.42	0.11	0.13	0.13
Ruminococcus CAG:382       0.10       0.03       0.03       0.03         Ruminococcus CAG:579       0.88       2.32       0.27       0.27         Saccharomonospora cyanea NA-134       0.13       0.04       0.04       0.04         Salinispora arenicola CNT849       0.86       0.23       0.26       0.26         Salinispora arenicola CNY234       0.61       0.17       0.19       0.19         Salinopella enterica SA-2       0.05       0.01       0.02       0.02         Salenonella enterica CFSAN001588       0.13       0.04       0.89       0.04         Selenomonas noxia ATCC 43541       0.06       0.02       0.02       0.02         Shewanella frigidimarina NCIMB 400       0.04       0.04       0.04       0.01         Shigella boydii 965-58       0.18       0.05       0.58       0.05         Shigella flexneri 1485-80       0.11       0.03       0.03       0.03         Simonsiella muelleri ATCC 29453       0.31       0.08       0.99       0.99         Sphingopyxis MC1       0.07       0.07       0.02       0.08         Streptococcus agalactiae GB00264       0.09       0.03       0.03       0.03         Streptococcus agalactiae MRI Z1-0	Ruminiclostridium thermocellum YS	0.55	0.15	1.98	0.17
Ruminococcus CAG:5790.882.320.270.27Saccharomonospora cyanea NA-1340.130.040.040.04Salinispora arenicola CNT8490.860.230.260.26Salinispora arenicola CNY2340.610.170.190.19Salinispora pacifica CNY3300.520.720.160.16Salmonella enterica SA-20.050.010.020.02Salmonella enterica CFSAN0015880.130.040.890.04Selenomonas noxia ATCC 435410.060.020.020.02Shewanella frigidimarina NCIMB 4000.040.040.040.01Shigella boydii 965-580.180.050.580.05Shigella flexneri 1485-800.110.030.030.03Shigella flexneri 2930-710.110.030.030.03Simonsiella muelleri ATCC 294530.310.080.090.09Sphingomonas melonis DAPP-PG 2240.480.130.150.15Shtaphylococcus agalactiae GB002640.090.050.030.03Streptococcus agalactiae MRI Z1-0220.380.900.120.12Streptococcus anginosus F02110.100.030.370.03	Ruminococcus CAG:382	0.10	0.03	0.03	0.03
Saccharomonospora cyanea NA-1340.130.040.040.04Salinispora arenicola CNT8490.860.230.260.26Salinispora arenicola CNY2340.610.170.190.19Salinispora pacifica CNY3300.520.720.160.16Salmonella enterica SA-20.050.010.020.02Salmonella enterica CFSAN0015880.130.040.890.04Selenomonas noxia ATCC 435410.060.020.020.02Shewanella frigidimarina NCIMB 4000.040.040.040.01Shigella boydii 965-580.180.050.580.05Shigella flexneri 1485-800.110.030.030.03Simonsiella muelleri ATCC 294530.310.080.090.09Sphingomonas melonis DAPP-PG 2240.480.130.150.15Sphingopyxis MC10.070.070.020.08Streptococcus agalactiae GB002640.090.090.030.03Streptococcus agalactiae MRI Z1-0220.380.900.120.12Streptococcus anginosus F02110.100.030.370.03	Ruminococcus CAG:579	0.88	2.32	0.27	0.27
Salinispora arenicola CNT8490.860.230.260.26Salinispora arenicola CNY2340.610.170.190.19Salinispora pacifica CNY3300.520.720.160.16Salmonella enterica SA-20.050.010.020.02Salmonella enterica CFSAN0015880.130.040.890.04Selenomonas noxia ATCC 435410.060.020.020.02Shewanella frigidimarina NCIMB 4000.040.040.040.01Shigella boydii 965-580.180.050.580.05Shigella flexneri 1485-800.110.030.030.03Shigella flexneri 2930-710.110.030.030.03Simonsiella muelleri ATCC 294530.310.080.090.09Sphingomonas melonis DAPP-PG 2240.480.130.150.15Sphingopyxis MC10.070.070.020.08Streptococcus sagalactiae GB002640.090.040.040.04Streptococcus agalactiae MRI Z1-0220.380.900.120.12Streptococcus anginosus F02110.100.030.370.03	Saccharomonospora cyanea NA-134	0.13	0.04	0.04	0.04
Salinispora arenicola CNY2340.610.170.190.19Salinispora pacifica CNY3300.520.720.160.16Salmonella enterica SA-20.050.010.020.02Salmonella enterica CFSAN0015880.130.040.890.04Selenomonas noxia ATCC 435410.060.020.020.02Shewanella frigidimarina NCIMB 4000.040.040.040.01Shigella boydii 965-580.180.050.580.05Shigella dysenteriae CDC 74-11120.201.730.500.45Shigella flexneri 1485-800.110.030.030.03Shigella flexneri 2930-710.110.080.090.09Sphingomonas melonis DAPP-PG 2240.480.130.150.15Sphingopyxis MC10.070.070.020.08Streptococcus agalactiae GB002640.090.090.030.03Streptococcus agalactiae MRI Z1-0220.380.900.120.12Streptococcus agalactiae MRI Z1-2020.380.900.120.12Streptococcus agalactiae MRI	Salinispora arenicola CNT849	0.86	0.23	0.26	0.26
Salinispora pacifica CNY3300.520.720.160.16Salmonella enterica SA-20.050.010.020.02Salmonella enterica CFSAN0015880.130.040.890.04Selenomonas noxia ATCC 435410.060.020.020.02Shewanella frigidimarina NCIMB 4000.040.040.040.01Shigella boydii 965-580.180.050.580.05Shigella dysenteriae CDC 74-11120.201.730.500.45Shigella flexneri 1485-800.110.030.030.03Shigella flexneri 2930-710.110.030.030.03Simonsiella muelleri ATCC 294530.310.080.090.09Sphingomonas melonis DAPP-PG 2240.480.130.150.15Sphingopyxis MC10.070.070.020.08Streptococcus agalactiae GB002640.090.050.030.03Streptococcus agalactiae MRI Z1-0220.380.900.120.12Streptococcus anginosus F02110.100.030.370.03	Salinispora arenicola CNY234	0.61	0.17	0.19	0.19
Salmonella enterica SA-20.050.010.020.02Salmonella enterica CFSAN0015880.130.040.890.04Selenomonas noxia ATCC 435410.060.020.020.02Shewanella frigidimarina NCIMB 4000.040.040.040.01Shigella boydii 965-580.180.050.580.05Shigella dysenteriae CDC 74-11120.201.730.500.45Shigella flexneri 1485-800.110.030.030.03Shigella flexneri 2930-710.110.030.030.03Simonsiella muelleri ATCC 294530.310.080.090.09Sphingomonas melonis DAPP-PG 2240.480.130.150.15Shighlylococcus hominis SK1190.090.030.030.03Streptococcus agalactiae GB002640.090.090.050.030.03Streptococcus agalactiae MRI Z1-0220.380.900.120.12Streptococcus anginosus F02110.100.030.370.03	Salinispora pacifica CNY330	0.52	0.72	0.16	0.16
Salmonella enterica CFSAN0015880.130.040.890.04Selenomonas noxia ATCC 435410.060.020.020.02Shewanella frigidimarina NCIMB 4000.040.040.040.01Shigella boydii 965-580.180.050.580.05Shigella dysenteriae CDC 74-11120.201.730.500.45Shigella flexneri 1485-800.110.030.030.03Shigella flexneri 2930-710.110.030.030.03Simonsiella muelleri ATCC 294530.310.080.090.09Sphingomonas melonis DAPP-PG 2240.480.130.150.15Staphylococcus hominis SK1190.090.090.030.030.03Streptococcus agalactiae GB002640.090.040.040.04Streptococcus agalactiae MRI Z1-0220.380.900.120.12Streptococcus anginosus F02110.100.030.370.03	Salmonella enterica SA-2	0.05	0.01	0.02	0.02
Selenomonas noxia ATCC 435410.060.020.020.02Shewanella frigidimarina NCIMB 4000.040.040.040.01Shigella boydii 965-580.180.050.580.05Shigella dysenteriae CDC 74-11120.201.730.500.45Shigella flexneri 1485-800.110.030.030.03Shigella flexneri 2930-710.110.030.030.03Simonsiella muelleri ATCC 294530.310.080.090.09Sphingomonas melonis DAPP-PG 2240.480.130.150.15Shigelylococcus hominis SK1190.090.030.030.03Streptococcus agalactiae GB002640.090.090.050.030.03Streptococcus agalactiae MRI Z1-0220.380.900.120.12Streptococcus anginosus F02110.100.030.370.03	Salmonella enterica CFSAN001588	0.13	0.04	0.89	0.04
Shewanella frigidimarina NCIMB 4000.040.040.040.01Shigella boydii 965-580.180.050.580.05Shigella dysenteriae CDC 74-11120.201.730.500.45Shigella flexneri 1485-800.110.030.030.03Shigella flexneri 2930-710.110.030.030.03Simonsiella muelleri ATCC 294530.310.080.090.09Sphingomonas melonis DAPP-PG 2240.480.130.150.15Sphingopyxis MC10.070.070.020.08Staphylococcus hominis SK1190.090.030.030.03Streptococcus agalactiae GB002640.090.140.040.04Streptococcus agalactiae MRI Z1-0220.380.900.120.12Streptococcus anginosus F02110.100.030.370.03	Selenomonas noxia ATCC 43541	0.06	0.02	0.02	0.02
Shigella boydii 965-580.180.050.580.05Shigella dysenteriae CDC 74-11120.201.730.500.45Shigella flexneri 1485-800.110.030.030.03Shigella flexneri 2930-710.110.030.030.03Simonsiella muelleri ATCC 294530.310.080.090.09Sphingomonas melonis DAPP-PG 2240.480.130.150.15Sphingopyxis MC10.070.070.020.08Staphylococcus hominis SK1190.090.090.030.03Streptococcus agalactiae GB002640.090.050.030.03Streptococcus agalactiae MRI Z1-0220.140.040.040.04Streptococcus anginosus F02110.100.030.370.03	Shewanella frigidimarina NCIMB 400	0.04	0.04	0.04	0.01
Shigella dysenteriae CDC 74-11120.201.730.500.45Shigella flexneri 1485-800.110.030.030.03Shigella flexneri 2930-710.110.030.030.03Simonsiella muelleri ATCC 294530.310.080.090.09Sphingomonas melonis DAPP-PG 2240.480.130.150.15Sphingopyxis MC10.070.070.020.08Staphylococcus hominis SK1190.090.030.030.03Streptococcus agalactiae GB002640.090.050.030.03Streptococcus agalactiae MRI Z1-0220.140.040.040.04Streptococcus agalactiae SP02110.100.030.370.03	Shigella boydii 965-58	0.18	0.05	0.58	0.05
Shigella flexneri 1485-800.110.030.030.03Shigella flexneri 2930-710.110.030.030.03Simonsiella muelleri ATCC 294530.310.080.090.09Sphingomonas melonis DAPP-PG 2240.480.130.150.15Sphingopyxis MC10.070.070.020.08Staphylococcus hominis SK1190.090.090.030.03Streptococcus agalactiae GB002640.090.050.030.03Streptococcus agalactiae MRI Z1-0220.140.040.040.04Streptococcus agalactiae SP02110.100.030.370.03	Shigella dysenteriae CDC 74-1112	0.20	1.73	0.50	0.45
Shigella flexneri 2930-710.110.030.030.03Simonsiella muelleri ATCC 294530.310.080.090.09Sphingomonas melonis DAPP-PG 2240.480.130.150.15Sphingopyxis MC10.070.070.020.08Staphylococcus hominis SK1190.090.030.030.03Streptococcus agalactiae GB002640.090.050.030.03Streptococcus agalactiae MRI Z1-0220.140.040.040.04Streptococcus agalactiae SP02110.100.030.370.03	Shigella flexneri 1485-80	0.11	0.03	0.03	0.03
Simonsiella muelleri ATCC 294530.310.080.090.09Sphingomonas melonis DAPP-PG 2240.480.130.150.15Sphingopyxis MC10.070.070.020.08Staphylococcus hominis SK1190.090.030.030.08Streptococcus agalactiae GB002640.090.050.030.03Streptococcus agalactiae MRI Z1-0220.140.040.040.04Streptococcus agalactiae MRI Z1-2020.380.900.120.12Streptococcus anginosus F02110.100.030.370.03	Shigella flexneri 2930-71	0.11	0.03	0.03	0.03
Sphingomonas melonis DAPP-PG 2240.480.130.150.15Sphingopyxis MC10.070.070.020.08Staphylococcus hominis SK1190.090.030.030.08Streptococcus agalactiae GB002640.090.050.030.03Streptococcus agalactiae MRI Z1-0220.140.040.040.04Streptococcus agalactiae MRI Z1-2020.380.900.120.12Streptococcus anginosus F02110.100.030.370.03	Simonsiella muelleri ATCC 29453	0.31	0.08	0.09	0.09
Sphingopyxis MC10.070.070.020.08Staphylococcus hominis SK1190.090.030.030.08Streptococcus agalactiae GB002640.090.050.030.03Streptococcus agalactiae MRI Z1-0220.140.040.040.04Streptococcus agalactiae MRI Z1-2020.380.900.120.12Streptococcus anginosus F02110.100.030.370.03	Sphingomonas melonis DAPP-PG 224	0.48	0.13	0.15	0.15
Staphylococcus hominis SK1190.090.030.030.08Streptococcus agalactiae GB002640.090.050.030.03Streptococcus agalactiae MRI Z1-0220.140.040.040.04Streptococcus agalactiae MRI Z1-2020.380.900.120.12Streptococcus anginosus F02110.100.030.370.03	Sphingopyxis MC1	0.07	0.07	0.02	0.08
Streptococcus agalactiae GB002640.090.050.030.03Streptococcus agalactiae MRI Z1-0220.140.040.040.04Streptococcus agalactiae MRI Z1-2020.380.900.120.12Streptococcus anginosus F02110.100.030.370.03	Staphylococcus hominis SK119	0.09	0.03	0.03	0.08
Streptococcus agalactiae MRI Z1-022       0.14       0.04       0.04         Streptococcus agalactiae MRI Z1-202       0.38       0.90       0.12       0.12         Streptococcus agalactiae S0211       0.10       0.03       0.37       0.03	Streptococcus agalactiae GB00264	0.09	0.05	0.03	0.03
Streptococcus agalactiae MRI Z1-202         0.38         0.90         0.12         0.12           Streptococcus anginosus F0211         0.10         0.03         0.37         0.03	Streptococcus agalactiae MRI Z1-022	0.14	0.04	0.04	0.04
Streptococcus anginosus F0211         0.10         0.03         0.37         0.03	Streptococcus agalactiae MRI Z1-202	0.38	0.90	0.12	0.12
	Streptococcus anginosus F0211	0.10	0.03	0.37	0.03

Streptococcus equi         0.37         0.10         1.54         0.11           Streptococcus efrus DSM 20646         0.15         0.04         0.05         0.05           Streptococcus iniae 9117         0.01         0.00         0.07         1.11         0.08           Streptococcus intermedius ATCC 27335         3.19         0.87         0.98         0.97           Streptococcus intermedius ATCC 27335         0.10         0.03         0.03         0.03           Streptococcus mutans KX23         0.62         0.17         0.19         0.19           Streptococcus salutans SM6         0.02         0.06         0.01         0.03           Streptococcus salutanis SX40         0.02         0.06         0.01         0.03           Streptococcus sobrinus TCI-367         1.65         0.45         0.50         0.50           Streptococcus Stl40         0.43         0.11         0.13         0.13         0.13           Streptococcus Stl40         0.43         0.12         0.13         0.13         0.13           Streptonyces acidiscables 84-104         0.22         0.26         0.07         0.07           Streptonyces sulphureus DSM 40104         0.13         0.11         0.42         0.12	Name	S1 (%)	S2 (%)	S3 (%)	S4 (%)
Streptococcus equi SzS31A1       0.31       0.08       0.09       0.27         Streptococcus ferus DSM 20646       0.15       0.04       0.05       0.05         Streptococcus gordoni CH       0.26       0.07       1.11       0.08         Streptococcus intermedius ATCC 27335       3.19       0.62       0.17       0.19       0.19         Streptococcus mutans SK6       0.10       0.03       0.03       0.03       0.03         Streptococcus sequiprices peudoportinus LQ 940-04       0.03       0.01       0.01       0.01       0.05         Streptococcus sobrinus DSM 20742/ATCC 33478       0.42       0.11       0.13       0.29         Streptococcus sobrinus TCI-367       1.65       0.45       0.50       0.50         Streptococcus St-P16       0.05       0.01       0.01       0.01         Streptococcus St-P16       0.05       0.01       0.01       0.13       0.13       0.13         Streptococcus St-P16       0.05       0.01       0.01       0.01       0.01       0.01       0.01       0.01       0.01       0.01       0.01       0.01       0.01       0.01       0.01       0.07       Streptococcus St-P16       0.05       0.01       0.01       0.02	Streptococcus equi	0.37	0.10	1.54	0.11
Streptococcus gordonii CH1       0.15       0.04       0.05       0.05         Streptococcus gordonii CH1       0.26       0.07       1.11       0.08         Streptococcus intermedius ATCC 27335       3.19       0.87       0.98       0.97         Streptococcus mutans KM23       0.62       0.67       0.19       0.19         Streptococcus salivarius 57.1       0.17       0.14       0.05       0.05         Streptococcus salivarius 57.1       0.17       0.14       0.05       0.55         Streptococcus salvirus 57.1       0.17       0.14       0.05       0.55         Streptococcus sobrinus TCI-367       1.66       0.45       0.50       55         Streptococcus sobrinus TCI-98       0.23       0.06       0.01       0.01         Streptococcus SK140       0.42       0.11       0.13       0.13       0.12       0.13       0.12         Streptococcus SK140       0.43       0.12       0.13       0.13       0.11       0.13       0.13       0.13       0.13       0.13       0.13       0.13       0.13       0.13       0.13       0.13       0.13       0.13       0.13       0.13       0.13       0.13       0.15       Streptococcus SK140       0	Streptococcus equi SzS31A1	0.31	0.08	0.09	0.27
Streptococcus gordonii CH1       0.26       0.07       1.11       0.08         Streptococcus intermedius ATCC 27335       3.19       0.67       0.98       0.97         Streptococcus mutans KK23       0.62       0.17       0.19       0.19         Streptococcus pseudoporcinus LQ 940-04       0.03       0.03       0.03       0.03         Streptococcus aspuints SX40       0.02       0.06       0.01       0.01         Streptococcus sobrinus DSM 20742/ATCC 33478       0.42       0.11       0.13       0.29         Streptococcus sobrinus TCI-367       1.66       0.45       0.01       0.01       0.01         Streptococcus sobrinus TCI-98       0.23       0.06       0.34       0.07       0.70         Streptococcus sobrinus TCI-98       0.42       0.11       0.13       0.13       0.13         Streptococcus subrinus TCI-98       0.42       0.11       0.13       0.11       0.16       0.11         Streptoxyces aidulachies 84-104       0.42       0.12       0.13       0.12       0.12       0.12       0.12       0.12       0.12       0.12       0.12       0.12       0.12       0.12       0.12       0.12       0.12       0.12       0.12       0.12       0.12 </td <td>Streptococcus ferus DSM 20646</td> <td>0.15</td> <td>0.04</td> <td>0.05</td> <td>0.05</td>	Streptococcus ferus DSM 20646	0.15	0.04	0.05	0.05
Streptococcus iniae 9117       0.01       0.00       0.01       0.09         Streptococcus intanes KK23       3.19       0.87       0.98       0.97         Streptococcus mutans SM6       0.10       0.03       0.01       0.01         Streptococcus salivarius 57.1       0.17       0.14       0.05       0.05         Streptococcus salivarius 57.1       0.17       0.14       0.05       0.05         Streptococcus sobrinus DSM 2074/ATCC 33478       0.42       0.11       0.13       0.29         Streptococcus sobrinus TCI-367       1.66       0.45       0.50       55         Streptococcus sobrinus TCI-98       0.23       0.06       0.01       0.01         Streptococcus SK140       0.42       0.11       0.13       0.13         Streptococcus sK140       0.42       0.11       0.12       0.13         Streptococcus skifus       0.41       0.22       0.66       0.07       0.07         Streptomyces acidiscables 84-104       0.22       0.26       0.07       0.07         Streptomyces sulplureus DSM 40104       0.13       0.11       0.27       0.08       0.08         Streptomyces vialacuaninger Tu 4113       0.27       0.08       0.08       0.08	Streptococcus gordonii CH1	0.26	0.07	1.11	0.08
Streptococcus intermedius ATCC 27335       3.19       0.67       0.98       0.97         Streptococcus mutans KK3       0.62       0.17       0.19       0.19         Streptococcus mutans SM6       0.10       0.03       0.03       0.03         Streptococcus saguinis SX40       0.02       0.06       0.01       0.03         Streptococcus sobrinus DSM 20742/ATCC 33478       0.42       0.11       0.13       0.29         Streptococcus sobrinus TCI-367       1.65       0.45       0.50       0.51         Streptococcus sobrinus TCI-367       1.65       0.45       0.50       0.01       0.01       0.01         Streptococcus sobrinus TCI-98       0.42       0.11       0.13       0.13       0.13         Streptococcus suis YE51       0.42       0.11       0.13       0.13         Streptomyces albulus CRC 11814       0.40       0.11       0.92       0.12         Streptomyces sulphureus DSM 40104       0.13       0.11       0.42       0.11       0.43       0.12         Streptomyces sulphureus DSM 40104       0.13       0.11       0.45       0.06       0.08         Succinatimonas hippi YIT 12066       0.10       0.33       0.44       0.03       0.44       0.03 <td>Streptococcus iniae 9117</td> <td>0.01</td> <td>0.00</td> <td>0.01</td> <td>0.09</td>	Streptococcus iniae 9117	0.01	0.00	0.01	0.09
Streptococcus mutans KK23       0.62       0.17       0.19       0.19         Streptococcus mutans SM6       0.10       0.03       0.03       0.03         Streptococcus salvarius SJ, I       0.17       0.14       0.05       0.05         Streptococcus salvarius SX40       0.02       0.01       0.01       0.03         Streptococcus sobrinus DSN 20742/ATCC 33478       0.42       0.11       0.13       0.29         Streptococcus sobrinus TCI-967       1.65       0.45       0.50       0.10       0.01       0.01         Streptococcus I-P16       0.05       0.01       0.01       0.01       0.11       0.13       0.13         Streptococcus Sulvinus TCI-98       0.42       0.11       0.13       0.13       0.13         Streptococcus Sulvinus TCC 198       0.42       0.12       0.13       0.13         Streptococcus Sulvinus CCR 11814       0.40       0.11       0.92       0.12         Streptomyces SUNProfe       0.17       0.05       0.15       0.05         Streptomyces VIQArce T1814       0.40       0.11       0.42       0.12       0.13       0.11         Streptomyces SUNPAres N40104       0.13       0.11       0.45       0.04       0.08	Streptococcus intermedius ATCC 27335	3.19	0.87	0.98	0.97
Streptococcus mutans SM6       0.10       0.03       0.03       0.03         Streptococcus salurius LQ 940-04       0.03       0.01       0.01         Streptococcus salurius S7.I       0.17       0.14       0.05       0.05         Streptococcus salurius SX40       0.02       0.06       0.01       0.03         Streptococcus sobrinus DCI-367       1.65       0.45       0.50       0.50         Streptococcus sobrinus TCI-367       1.65       0.43       0.01       0.01         Streptococcus sobrinus TCI-98       0.23       0.06       0.34       0.07         Streptococcus SX140       0.42       0.11       0.13       0.13         Streptococcus SX140       0.42       0.11       0.13       0.13         Streptomyces alculus CCR 11814       0.40       0.11       0.92       0.12         Streptomyces sulphureu DSM 40104       0.13       0.15       0.05         Streptomyces sulphureu DSM 40104       0.13       0.11       0.44       0.03         Sulfolobus islandicus REY15A       0.10       0.03       0.44       0.03         Synechocytis PCC 6803       0.42       0.12       0.13       0.11         Synechocytis PCC 7509       0.16       0.16 <td>Streptococcus mutans KK23</td> <td>0.62</td> <td>0.17</td> <td>0.19</td> <td>0.19</td>	Streptococcus mutans KK23	0.62	0.17	0.19	0.19
Streptococcus pseudoporcinus LQ 940-04       0.03       0.01       0.01       0.01         Streptococcus sanguinis SK340       0.02       0.06       0.05         Streptococcus sobrinus DSN 20742/ATCC 33478       0.42       0.11       0.13       0.29         Streptococcus sobrinus TCI-367       1.65       0.46       0.01       0.01         Streptococcus sobrinus TCI-367       1.65       0.42       0.11       0.13       0.12         Streptococcus sobrinus TCI-98       0.23       0.06       0.34       0.07       0.01       0.01         Streptococcus sobrinus TCI-98       0.42       0.11       0.13       0.13       0.13         Streptococcus signinus TCI-98       0.42       0.11       0.13       0.13       0.13         Streptomyces acidiscables 84-104       0.22       0.26       0.07       0.07         Streptomyces pristinaespiralis ATCC 25486       0.11       0.12       0.13       0.15       0.05         Streptomyces violaceusniger Tu 413       0.27       0.08       0.08       0.08       0.08         Succinatimonas hippei YIT 12066       0.10       0.03       0.33       0.08       0.01         Synechocystis PCC 6803       0.42       0.12       0.13       0.	Streptococcus mutans SM6	0.10	0.03	0.03	0.03
Streptococcus salivarius 57.I       0.17       0.14       0.05       0.05         Streptococcus sanguins SX40       0.02       0.06       0.01       0.03         Streptococcus sobrinus DSM 20742/ATCC 33478       0.42       0.11       0.13       0.29         Streptococcus sobrinus TCI-367       1.65       0.45       0.50       0.50         Streptococcus sobrinus TCI-98       0.23       0.06       0.34       0.01       0.01         Streptococcus SK140       0.43       0.12       0.13       0.13       0.13         Streptocccus SK140       0.42       0.11       0.13       0.13       0.13         Streptomyces albulus CCR 11814       0.40       0.11       0.27       0.03       0.09         Streptomyces SUR766       0.17       0.05       0.15       0.05         Streptomyces violaceusinger Tu 4113       0.27       0.08       0.08       0.08         Sulfalobus islandicus REVISA       0.10       0.03       0.03       0.03       0.03         Synechocystis PCC 7509       0.04       0.05       0.01       0.01       0.01         Synechocystis PCC 7509       0.04       0.05       0.05       0.05       0.05         Thermotoga elfin NBRC 107921<	Streptococcus pseudoporcinus LQ 940-04	0.03	0.01	0.01	0.01
Streptococcus sanguinis SX340       0.02       0.06       0.01       0.03         Streptococcus sobrinus DIM 20742/ATCC 33478       0.42       0.11       0.13       0.29         Streptococcus sobrinus TCI-98       0.23       0.06       0.34       0.07         Streptococcus sobrinus TCI-98       0.23       0.06       0.34       0.07         Streptococcus Sti40       0.43       0.12       0.13       0.13         Streptococcus suis YB51       0.42       0.11       0.13       0.13         Streptomyces albulus CCRC 11814       0.40       0.21       0.70       0.07         Streptomyces Streptomyces NUP766       0.17       0.05       0.15       0.05         Streptomyces violaceuniger Tu 4113       0.27       0.08       0.08       0.08         Succinatinonas hippei YIT 12066       0.10       0.03       0.44       0.03         Synechoccystis PCC 7509       0.04       0.05       0.01       0.01         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.64	Streptococcus salivarius 57.I	0.17	0.14	0.05	0.05
Streptococcus sobrinus DSN 20742/ATCC 33478       0.42       0.11       0.13       0.29         Streptococcus sobrinus TCI-367       1.65       0.45       0.50       0.07         Streptococcus sobrinus TCI-98       0.23       0.06       0.34       0.07         Streptococcus solvanus TCI-98       0.43       0.11       0.13       0.13         Streptococcus suis YB51       0.42       0.11       0.13       0.13         Streptomyces acidiscabies 84-104       0.22       0.26       0.07       0.07         Streptomyces acidiscabies 84-104       0.40       0.11       0.92       0.12         Streptomyces pristinaespiralis ATCC 25486       0.11       0.45       0.03       0.09         Streptomyces sulphureus DSM 40104       0.13       0.11       0.45       0.04         Streptomyces sulphureus DSM 40104       0.13       0.11       0.45       0.04         Succinationas hipper YT 12066       0.10       0.03       0.03       0.08         Succinationas hipper YT 12066       0.10       0.03       0.44       0.03         Synechocystis PCC 7509       0.04       0.05       0.01       0.01         Synechocystis PCC 7509       0.04       0.04       0.04       0.04 <td>Streptococcus sanguinis SK340</td> <td>0.02</td> <td>0.06</td> <td>0.01</td> <td>0.03</td>	Streptococcus sanguinis SK340	0.02	0.06	0.01	0.03
Streptococcus sobrinus TCI-967       1.65       0.45       0.50       0.50         Streptococcus sobrinus TCI-98       0.23       0.06       0.34       0.07         Streptococcus SK140       0.43       0.12       0.13       0.13         Streptococcus SK140       0.42       0.12       0.13       0.13         Streptococcus suis YB51       0.42       0.26       0.07       0.07         Streptomyces albulus CCRC 11814       0.40       0.11       0.92       0.12         Streptomyces Sulphureus DSM 40104       0.13       0.11       0.45       0.04         Streptomyces violaceusniger Tu 4113       0.27       0.08       0.08       0.08         Sulfolobus islandicus REY15A       0.10       0.03       0.04       0.01       0.30       0.04         Synechocystis PCC 6803       0.04       0.01       0.30       0.04       0.01       0.30       0.04         Thermosphera aggregans DSM 11486       0.69       0.04       0.04       0.04       0.04       0.04         Thermosphera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermosphera aggregans DSM 11486       0.69       0.44       0.10       0.04       0.04       0.04	Streptococcus sobrinus DSM 20742/ATCC 33478	0.42	0.11	0.13	0.29
Streptococcus sobrinus TCI-98       0.23       0.06       0.34       0.07         Streptococcus SK140       0.05       0.01       0.01       0.01         Streptococcus suis YB51       0.42       0.11       0.13       0.13         Streptomyces acidiscabies 84-104       0.22       0.26       0.07       0.07         Streptomyces acidiscabies 84-104       0.42       0.11       0.92       0.12         Streptomyces acidiscabies 84-104       0.40       0.11       0.92       0.09         Streptomyces pristinaespiralis ATCC 25486       0.11       0.27       0.03       0.09         Streptomyces violaceusniger Tu 4113       0.27       0.08       0.08       0.08         Succinationas hippei VIT 12066       0.10       0.03       0.44       0.03         Synechocystis PCC 7836       0.42       0.12       0.13       0.11         Synechocystis PCC 7509       0.04       0.05       0.01       0.03       0.04         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermotoga elfii NBRC 107921       0.16       0.04       0.24       0.91	Streptococcus sobrinus TCI-367	1.65	0.45	0.50	0.50
Streptococcus I-P16       0.05       0.01       0.01       0.01         Streptococcus SX140       0.43       0.12       0.13       0.13         Streptococcus suis YB51       0.42       0.11       0.13       0.13         Streptomyces acidiscabies 84-104       0.40       0.11       0.92       0.12         Streptomyces pristinaespiralis ATCC 25486       0.11       0.47       0.03       0.09         Streptomyces cNQ766       0.17       0.05       0.15       0.05         Streptomyces violaceusniger Tu 4113       0.27       0.08       0.08       0.08         Succinatimonas hippei YIT 12066       0.10       0.03       0.44       0.03         Synechocycus PCC 7336       0.42       0.12       0.13       0.11         Synechocystis PCC 6803       0.04       0.05       0.01       0.03         Synechocystis PCC 7509       0.04       0.05       0.01       0.04         Thauera linalcolentis 47Lo1/DSM 12138       0.28       0.08       0.08       0.08         Thermospaera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermospaera aggregans DSM 11486       0.36       0.10       0.11       0.11         Thermotoga EMP	Streptococcus sobrinus TCI-98	0.23	0.06	0.34	0.07
Streptococcus SK140       0.43       0.12       0.13       0.13         Streptococcus suis YB51       0.42       0.11       0.13       0.13         Streptomyces alcidiscables 84-104       0.22       0.26       0.07       0.07         Streptomyces alcidiscables 84-104       0.40       0.11       0.92       0.12         Streptomyces pristinaespiralis ATCC 25486       0.11       0.27       0.03       0.09         Streptomyces sulphureus DSM 40104       0.13       0.11       0.45       0.04         Streptomyces vialaceusniger Tu 4113       0.27       0.08       0.08       0.08         Succinatimonas hippei YIT 12066       0.10       0.03       0.04       0.03       0.04         Synechocystis PCC 7366       0.42       0.12       0.13       0.11         Synechocystis PCC 7509       0.04       0.05       0.01       0.01         Thermotoga elfii NBRC 107921       0.16       0.04       0.04       0.04         Thermotoga elfii NBRC 107921       0.16       0.04       0.24       0.12       0.13         Thermotoga elfii NBRC 107921       0.16       0.46       0.12       0.13         Thermotoga elfii NBRC 107921       0.16       0.04       0.24	Streptococcus I-P16	0.05	0.01	0.01	0.01
Streptococcus suis YB51       0.42       0.11       0.13       0.13         Streptomyces acidiscabies 84-104       0.22       0.26       0.07       0.07         Streptomyces albulus CCRC 11814       0.40       0.11       0.92       0.12         Streptomyces pristinaespiralis ATCC 25486       0.11       0.27       0.03       0.09         Streptomyces violaccusniger Tu 4113       0.27       0.08       0.08       0.08         Succinatimonas hippei YIT 12066       0.10       0.03       0.04       0.03         Synechocycstis PCC 7336       0.42       0.12       0.13       0.11         Synechocystis PCC 7509       0.04       0.05       0.01       0.30       0.04         Thermococcus onnurineus NA1       0.15       0.18       0.09       0.04         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.164         Thermospaera aggregans DSM 11486       0.69       0.74       0.21       1.164         Thermotoga EMP       0.51       0.14       0.16       3.02       0.11       0.11       0.11         Thermotoga EMP       0.51       0.14       0.16       3.02       0.66       0.25       0.15         Theolakalivibrio ALJ	Streptococcus SK140	0.43	0.12	0.13	0.13
Streptomyces acidiscabies 84-104       0.22       0.26       0.07       0.07         Streptomyces albulus CCRC 11814       0.40       0.11       0.92       0.12         Streptomyces pristinaespiralis ATCC 25486       0.11       0.27       0.03       0.09         Streptomyces Sulphureus DSM 40104       0.13       0.11       0.45       0.04         Streptomyces violaceusinger Tu 4113       0.27       0.08       0.08       0.03         Sulfolobus islandicus REY15A       0.10       0.03       0.04       0.03       0.04         Synechococcus PCC 7336       0.42       0.12       0.13       0.11       0.30       0.01         Synechocystis PCC 6803       0.04       0.01       0.30       0.04       0.01       0.30       0.04         Thearea linaloolentis 47Lol/DSM 12138       0.28       0.08       0.08       0.08       0.08         Thermotoga elfii NBRC 107921       0.16       0.04       0.04       0.04       0.04       0.11       0.11       0.11       0.11         Theomotoga EMP       0.51       0.14       0.16       0.02       0.05       0.05       0.05         Theomotoga ELYS_UFI       0.17       0.55       0.05       0.16       0.10	Streptococcus suis YB51	0.42	0.11	0.13	0.13
Streptomyces albulus CCRC 11814       0.40       0.11       0.92       0.12         Streptomyces pristinaespiralis ATCC 25486       0.11       0.27       0.03       0.09         Streptomyces Sulphureus DSM 40104       0.13       0.11       0.45       0.04         Streptomyces violaceusniger Tu 4113       0.27       0.08       0.08       0.08         Succinatimonas hippei YIT 12066       0.10       0.03       0.03       0.08         Sulfolobus islandicus REY15A       0.10       0.03       0.44       0.03         Synechococcus PCC 7336       0.42       0.12       0.13       0.11         Synechocystis PCC 6803       0.04       0.01       0.30       0.04         Synechocystis PCC 7509       0.04       0.05       0.01       0.01         Thermosphaera lagregans DSM 112138       0.28       0.08       0.08       0.08         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermosphaera aggregans DSM 11486       0.60       0.60       0.65       0.05	Streptomyces acidiscabies 84-104	0.22	0.26	0.07	0.07
Streptomyces pristinaespiralis ATCC 25486       0.11       0.27       0.03       0.09         Streptomyces CNQ766       0.17       0.05       0.15       0.05         Streptomyces sulphureus DSM 40104       0.13       0.11       0.45       0.04         Streptomyces violaceusniger Tu 4113       0.27       0.08       0.08       0.08         Succinatimonas hippei YIT 12066       0.10       0.03       0.04       0.03       0.03       0.08         Sunchococcus PCC 7336       0.42       0.12       0.13       0.13       0.11       0.30       0.01         Synechocystis PCC 7509       0.04       0.05       0.01       0.00       0.04       0.04       0.04         Thermococcus onnurineus NA1       0.15       0.18       0.09       0.04       0.04       0.04       0.04         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermotoga elfii NBRC 107921       0.16       0.04       0.04       0.04         Thermotoga EMP       0.51       0.14       0.16       3.02         Thermus CCE_US3_UF1       0.16       0.04       0.22       0.51         Thioalkalivibrio ALJ12       0.81       0.22       0.65	Streptomyces albulus CCRC 11814	0.40	0.11	0.92	0.12
Streptomyces CNQ766       0.17       0.05       0.15       0.05         Streptomyces sulphureus DSM 40104       0.13       0.11       0.45       0.04         Streptomyces violaceusniger Tu 4113       0.27       0.08       0.08       0.08         Succinatimonas hippei YIT 12066       0.10       0.03       0.04       0.03         Sulfolobus islandicus REY15A       0.10       0.03       0.44       0.03         Synechococcus PCC 7336       0.42       0.12       0.13       0.11         Synechocystis PCC 6803       0.04       0.05       0.01       0.01         Synechocystis PCC 7509       0.04       0.05       0.01       0.01         Thauera linaloolentis 47Lol/DSM 12138       0.28       0.08       0.08       0.08         Thermoccus onnurineus NA1       0.15       0.18       0.09       0.04         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermotoga EMP       0.51       0.14       0.16       3.02         Thioalkalivibrio ALJ0       0.60       2.66       0.19       0.18         Thioalkalivibrio ALJ10       0.60       2.66       0.19       0.18         Thioalkalivibrio ALJ24       0.4	Streptomyces pristinaespiralis ATCC 25486	0.11	0.27	0.03	0.09
Streptomyces sulphureus DSM 40104       0.13       0.11       0.45       0.04         Streptomyces violaceusniger Tu 4113       0.27       0.08       0.08       0.08         Succinatimonas hippei YIT 12066       0.10       0.03       0.03       0.03         Sulfolobus islandicus REY15A       0.10       0.03       0.44       0.03         Synechococcus PCC 7336       0.42       0.12       0.13       0.11         Synechocystis PCC 6803       0.04       0.01       0.30       0.01         Synechocystis PCC 7509       0.04       0.05       0.01       0.01         Thauera linaloolentis 47Lo1/DSM 12138       0.28       0.08       0.08       0.08         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermotoga elfii NBRC 107921       0.16       0.04       0.24       0.91         Thermus CCE_US3_UF1       0.17       0.05       0.05       0.05         Thioalkalivibrio ALL20       0.38       0.61       0.12       0.11         Thioalkalivibrio ALJ10       0.60       2.66       0.25       0.15         Thioalkalivibrio ALJ5       0.10       0.03       0.03       0.03         Thioalkalivibrio ALJ5	Streptomyces CNQ766	0.17	0.05	0.15	0.05
Streptomyces violaceusniger Tu 4113       0.27       0.08       0.08       0.08         Succinatimonas hippei YIT 12066       0.10       0.03       0.03       0.08         Sulfolobus islandicus REY15A       0.10       0.03       0.44       0.03         Synechococcus PCC 7336       0.42       0.12       0.13       0.13         Synechocystis PCC 6803       0.04       0.01       0.30       0.04         Synechocystis PCC 7509       0.04       0.05       0.01       0.01         Thauera linaloolentis 47L01/DSM 12138       0.28       0.08       0.08       0.08         Thermococcus onnurineus NA1       0.15       0.18       0.09       0.04         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermotoga EMP       0.51       0.14       0.16       3.02         Thermotoga EMP       0.51       0.14       0.16       3.02         Thioalkalivibrio ALE20       0.38       0.61       0.12       0.11         Thioalkalivibrio ALJ10       0.60       2.66       0.19       0.18         Thioalkalivibrio ALJ24       0.48       3.07       0.15       0.15         Thioalkalivibrio ALJ9       0.10       <	Streptomyces sulphureus DSM 40104	0.13	0.11	0.45	0.04
Succinatimonas hippei YIT 12066       0.10       0.03       0.03       0.08         Sulfolobus islandicus REY15A       0.10       0.03       0.44       0.03         Synechcoccus PCC 7336       0.42       0.12       0.13       0.13         Synechcocystis PCC 6803       0.04       0.01       0.30       0.01         Synechcystis PCC 7509       0.04       0.05       0.01       0.01         Thauera linaloolentis 47Lo1/DSM 12138       0.28       0.08       0.08       0.08         Thermococcus onnurineus NA1       0.15       0.18       0.09       0.04         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermotoga elfii NBRC 107921       0.16       0.04       0.24       0.91         Thermotoga EMP       0.51       0.14       0.16       3.02         Thrioalkalivibrio ALE20       0.38       0.61       0.12       0.11         Thioalkalivibrio ALJ10       0.60       2.66       0.19       0.18         Thioalkalivibrio ALJ24       0.48       3.07       0.15       0.15         Thioalkalivibrio ALJ5       0.10       0.03       0.28       0.47         Thioalkalivibrio ALJ5       0.10       <	Streptomyces violaceusniger Tu 4113	0.27	0.08	0.08	0.08
Sulfolobus islandicus REY15A       0.10       0.03       0.44       0.03         Synechococcus PCC 7336       0.42       0.12       0.13       0.13         Synechocystis PCC 6803       0.04       0.01       0.30       0.01         Synechocystis PCC 7509       0.04       0.05       0.01       0.01         Thauera linaloolentis 47Lol/DSM 12138       0.28       0.08       0.08       0.08         Thermococcus onnurineus NA1       0.15       0.18       0.09       0.04         Thermosphaera aggregans DSM 11486       0.69       0.74       0.21       1.64         Thermotoga elfii NBRC 107921       0.16       0.04       0.24       0.91         Thermotoga EMP       0.51       0.14       0.16       3.02         Thioalkalivibrio ALL20       0.38       0.61       0.12       0.11         Thioalkalivibrio ALJ10       0.60       2.66       0.19       0.18         Thioalkalivibrio ALJ24       0.48       3.07       0.15       0.15         Thioalkalivibrio ALJ5       0.10       0.03       0.28       0.47         Thioalkalivibrio ALJ24       0.48       3.07       0.15       0.15         Thioalkalivibrio ALJ5       0.10       0.03 </td <td>Succinatimonas hippei YIT 12066</td> <td>0.10</td> <td>0.03</td> <td>0.03</td> <td>0.08</td>	Succinatimonas hippei YIT 12066	0.10	0.03	0.03	0.08
Synechococcus PCC 73360.420.120.130.13Synechocystis PCC 68030.040.010.300.01Synechocystis PCC 75090.040.050.010.01Thauera linaloolentis 47Lol/DSM 121380.280.080.080.08Thermococcus onnurineus NA10.150.180.090.04Thermosphaera aggregans DSM 114860.690.740.211.64Thermotoga elfii NBRC 1079210.160.040.140.16Thermotoga EMP0.510.140.163.02Thrialkalivibrio AKL60.360.100.110.11Thioalkalivibrio ALJ200.380.610.120.11Thioalkalivibrio ALJ120.810.220.650.25Thioalkalivibrio ALJ240.483.070.150.15Thioalkalivibrio ALJ90.100.030.280.47Thioalkalivibrio ALJ90.160.040.050.05Ureaplasma urealyticum ATCC 278140.320.091.240.10Variuvorax paradoxus S1100.080.020.110.07Vibrio cholerae HC-50A20.160.040.050.05	Sulfolobus islandicus REY15A	0.10	0.03	0.44	0.03
Synechocystis PCC 68030.040.010.300.01Synechocystis PCC 75090.040.050.010.01Thauera linaloolentis 47Lol/DSM 121380.280.080.080.08Thermococcus onnurineus NA10.150.180.090.04Thermoplasmatales archaeon I-plasma0.130.040.040.04Thermosphaera aggregans DSM 114860.690.740.211.64Thermotoga elfii NBRC 1079210.160.040.240.91Thermotoga EMP0.510.140.163.02Throalkalivibrio AKL60.360.100.110.11Thioalkalivibrio AL200.380.610.120.11Thioalkalivibrio ALJ100.602.660.190.18Thioalkalivibrio ALJ240.483.070.150.15Thioalkalivibrio ALJ90.100.030.280.47Thioalkalivibrio ALJ90.100.030.030.03Tyzzerella nexilis DSM 17870.160.040.050.05uncultured archaeon A07HR600.670.182.135.67Ureaplasma urealyticum ATCC 278140.320.091.240.10Verucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	Synechococcus PCC 7336	0.42	0.12	0.13	0.13
Synechocystis PCC 75090.040.050.010.01Thauera linaloolentis 47Lo1/DSM 121380.280.080.080.08Thermococcus onnurineus NA10.150.180.090.04Thermoplasmatales archaeon I-plasma0.130.040.040.04Thermosphaera aggregans DSM 114860.690.740.211.64Thermotoga elfii NBRC 1079210.160.040.240.91Thermotoga EMP0.510.140.163.02Thermus CCE_US3_UF10.170.050.050.05Thioalkalivibrio ALE200.380.610.120.11Thioalkalivibrio ALJ120.810.220.650.25Thioalkalivibrio ALJ240.483.070.150.15Thioalkalivibrio ALJ50.100.030.280.47Thioalkalivibrio ALJ90.160.040.050.05Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45	Synechocystis PCC 6803	0.04	0.01	0.30	0.01
Thauera linaloolentis 47Lol/DSM 121380.280.080.080.08Thermococcus onnurineus NA10.150.180.090.04Thermoplasmatales archaeon I-plasma0.130.040.040.04Thermosphaera aggregans DSM 114860.690.740.211.64Thermotoga elfii NBRC 1079210.160.040.240.91Thermotoga EMP0.510.140.163.02Thermus CCB_US3_UF10.170.050.050.05Thioalkalivibrio AKL60.360.100.110.11Thioalkalivibrio AL200.380.610.120.11Thioalkalivibrio ALJ100.602.660.190.18Thioalkalivibrio ALJ240.483.070.150.15Thioalkalivibrio ALJ50.100.030.030.03Tyzzerella nexilis DSM 17870.160.040.050.05uncultured archaeon A07HR600.670.182.135.67Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	Synechocystis PCC 7509	0.04	0.05	0.01	0.01
Thermococcus onnurineus NA10.150.180.090.04Thermoplasmatales archaeon I-plasma0.130.040.040.04Thermosphaera aggregans DSM 114860.690.740.211.64Thermotoga elfii NBRC 1079210.160.040.240.91Thermotoga EMP0.510.140.163.02Thermus CCB_US3_UF10.170.050.050.05Thioalkalivibrio AKL60.360.100.110.11Thioalkalivibrio AL200.380.610.120.11Thioalkalivibrio ALJ100.602.660.190.18Thioalkalivibrio ALJ240.483.070.150.15Thioalkalivibrio ALJ50.100.030.280.47Thioalkalivibrio ALJ90.160.040.050.05Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45	Thauera linaloolentis 47Lol/DSM 12138	0.28	0.08	0.08	0.08
Thermoplasmatales archaeon I-plasma0.130.040.040.04Thermosphaera aggregans DSM 114860.690.740.211.64Thermotoga elfii NBRC 1079210.160.040.240.91Thermotoga EMP0.510.140.163.02Thermus CCB_US3_UF10.170.050.050.05Thioalkalivibrio AKL60.360.100.110.11Thioalkalivibrio AL200.380.610.120.11Thioalkalivibrio ALJ100.602.660.190.18Thioalkalivibrio ALJ120.483.070.150.15Thioalkalivibrio ALJ240.483.070.150.15Thioalkalivibrio ALJ50.100.030.030.03Thioalkalivibrio ALJ90.100.030.030.03Thioalkalivibrio ALJ90.160.040.050.05Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Vibrio cholerae HC-50A20.160.040.050.05	Thermococcus onnurineus NA1	0.15	0.18	0.09	0.04
Thermosphaera aggregans DSM 114860.690.740.211.64Thermotoga elfii NBRC 1079210.160.040.240.91Thermotoga EMP0.510.140.163.02Thermus CCB_US3_UF10.170.050.050.05Thioalkalivibrio AKL60.360.100.110.11Thioalkalivibrio AL200.380.610.120.11Thioalkalivibrio ALJ100.602.660.190.18Thioalkalivibrio ALJ240.810.220.650.25Thioalkalivibrio ALJ50.100.030.280.47Thioalkalivibrio ALJ90.160.040.050.05Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Vibrio cholerae HC-50A20.160.040.050.05	Thermoplasmatales archaeon I-plasma	0.13	0.04	0.04	0.04
Thermotoga elfii NBRC 1079210.160.040.240.91Thermotoga EMP0.510.140.163.02Thermus CCB_US3_UF10.170.050.050.05Thioalkalivibrio AKL60.360.100.110.11Thioalkalivibrio ALE200.380.610.120.11Thioalkalivibrio ALJ100.602.660.190.18Thioalkalivibrio ALJ120.810.220.650.25Thioalkalivibrio ALJ240.483.070.150.15Thioalkalivibrio ALJ50.100.030.280.47Thioalkalivibrio ALJ90.160.040.050.05uncultured archaeon A07HR600.670.182.135.67Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	Thermosphaera aggregans DSM 11486	0.69	0.74	0.21	1.64
Thermotoga EMP0.510.140.163.02Thermus CCB_US3_UF10.170.050.050.05Thioalkalivibrio AKL60.360.100.110.11Thioalkalivibrio ALE200.380.610.120.11Thioalkalivibrio ALJ100.602.660.190.18Thioalkalivibrio ALJ120.810.220.650.25Thioalkalivibrio ALJ240.483.070.150.15Thioalkalivibrio ALJ50.100.030.280.47Thioalkalivibrio ALJ90.100.030.030.03Tyzzerella nexilis DSM 17870.160.040.050.05Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	Thermotoga elfii NBRC 107921	0.16	0.04	0.24	0.91
Thermus CCB_US3_UF10.170.050.050.05Thioalkalivibrio AKL60.360.100.110.11Thioalkalivibrio ALE200.380.610.120.11Thioalkalivibrio ALJ100.602.660.190.18Thioalkalivibrio ALJ120.810.220.650.25Thioalkalivibrio ALJ240.483.070.150.15Thioalkalivibrio ALJ50.100.030.280.47Thioalkalivibrio ALJ90.100.030.030.03Tyzzerella nexilis DSM 17870.160.040.050.05Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Vibrio cholerae HC-50A20.160.040.050.05	Thermotoga EMP	0.51	0.14	0.16	3.02
Thioalkalivibrio AKL60.360.100.110.11Thioalkalivibrio AL200.380.610.120.11Thioalkalivibrio ALJ100.602.660.190.18Thioalkalivibrio ALJ120.810.220.650.25Thioalkalivibrio ALJ240.483.070.150.15Thioalkalivibrio ALJ50.100.030.280.47Thioalkalivibrio ALJ90.100.030.030.03Tyzzerella nexilis DSM 17870.160.040.050.05uncultured archaeon A07HR600.670.182.135.67Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	Thermus CCB_US3_UF1	0.17	0.05	0.05	0.05
Thioalkalivibrio ALE200.380.610.120.11Thioalkalivibrio ALJ100.602.660.190.18Thioalkalivibrio ALJ120.810.220.650.25Thioalkalivibrio ALJ240.483.070.150.15Thioalkalivibrio ALJ50.100.030.280.47Thioalkalivibrio ALJ90.100.030.030.03Tyzzerella nexilis DSM 17870.160.040.050.05uncultured archaeon A07HR600.670.182.135.67Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	Thioalkalivibrio AKL6	0.36	0.10	0.11	0.11
Thioalkalivibrio ALJ100.602.660.190.18Thioalkalivibrio ALJ120.810.220.650.25Thioalkalivibrio ALJ240.483.070.150.15Thioalkalivibrio ALJ50.100.030.280.47Thioalkalivibrio ALJ90.100.030.030.03Tyzzerella nexilis DSM 17870.160.040.050.05uncultured archaeon A07HR600.670.182.135.67Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	Thioalkalivibrio ALE20	0.38	0.61	0.12	0.11
Thioalkalivibrio ALJ120.810.220.650.25Thioalkalivibrio ALJ240.483.070.150.15Thioalkalivibrio ALJ50.100.030.280.47Thioalkalivibrio ALJ90.100.030.030.03Tyzzerella nexilis DSM 17870.160.040.050.05uncultured archaeon A07HR600.670.182.135.67Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	Thioalkalivibrio ALJ10	0.60	2.66	0.19	0.18
Thioalkalivibrio ALJ240.483.070.150.15Thioalkalivibrio ALJ50.100.030.280.47Thioalkalivibrio ALJ90.100.030.030.03Tyzzerella nexilis DSM 17870.160.040.050.05uncultured archaeon A07HR600.670.182.135.67Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	Thioalkalivibrio ALJ12	0.81	0.22	0.65	0.25
Thioalkalivibrio ALJ50.100.030.280.47Thioalkalivibrio ALJ90.100.030.030.03Tyzzerella nexilis DSM 17870.160.040.050.05uncultured archaeon A07HR600.670.182.135.67Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	Thioalkalivibrio ALJ24	0.48	3.07	0.15	0.15
Thioalkalivibrio ALJ90.100.030.030.03Tyzzerella nexilis DSM 17870.160.040.050.05uncultured archaeon A07HR600.670.182.135.67Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	Thioalkalivibrio ALJ5	0.10	0.03	0.28	0.47
Tyzzerella nexilis DSM 17870.160.040.050.05uncultured archaeon A07HR600.670.182.135.67Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	Thioalkalivibrio ALJ9	0.10	0.03	0.03	0.03
uncultured archaeon A07HR600.670.182.135.67Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	Tyzzerella nexilis DSM 1787	0.16	0.04	0.05	0.05
Ureaplasma urealyticum ATCC 278140.320.091.240.10Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	uncultured archaeon A07HR60	0.67	0.18	2.13	5.67
Variovorax paradoxus S1100.080.020.110.07Verrucomicrobium 3C1.487.162.750.45Vibrio cholerae HC-50A20.160.040.050.05	Ureaplasma urealyticum ATCC 27814	0.32	0.09	1.24	0.10
Verrucomicrobium 3C         1.48         7.16         2.75         0.45           Vibrio cholerae HC-50A2         0.16         0.04         0.05         0.05	Variovorax paradoxus S110	0.08	0.02	0.11	0.07
Vibrio cholerae HC-50A2         0.16         0.04         0.05         0.05	Verrucomicrobium 3C	1.48	7.16	2.75	0.45
	Vibrio cholerae HC-50A2	0.16	0.04	0.05	0.05

Name	S1 (%)	S2 (%)	S3 (%)	S4 (%)
Vibrio cholerae HE39	0.08	0.02	0.02	0.02
Vibrio cholerae O1 str. 2009V-1085	0.03	0.01	0.08	0.01
Vibrio crassostreae 9ZC88	0.18	0.05	0.05	0.05
Vibrio gazogenes ATCC 43941	0.96	2.34	0.29	0.29
Vibrio nigripulchritudo ENn2	0.71	0.20	0.22	0.22
Vibrio nigripulchritudo SFn135	0.22	0.06	1.80	0.07
Vibrio nigripulchritudo SOn1	0.46	0.13	0.14	0.14
Weissella koreensis KACC 15510	0.25	0.38	0.25	0.08
Wolbachia endosymbiont JHB	0.39	0.11	0.12	0.12
Xanthomonas axonopodis IBSBF 614	0.06	0.02	0.10	0.02
Xanthomonas axonopodis UA306	0.15	0.04	0.05	0.05
Xanthomonas campestris NCPPB 2005	0.17	0.05	0.05	0.05
Xanthomonas oryzae BLS256	0.18	0.05	0.06	0.06
Xanthomonas SHU166	0.13	0.04	0.04	0.04
Xylella fastidiosa 32	0.18	0.23	0.05	0.05
Yersinia frederiksenii ATCC 33641	0.22	1.08	0.07	1.20
Yersinia pseudotuberculosis B-6863	0.22	0.06	0.07	0.07
Yersinia pseudotuberculosis B-6864	0.12	1.02	0.13	0.19



Supplementary Figure 1: Genome enrichment for 400 genomes in the three-fold cross-validation. For each genome, we measured the sensitivity, the percentage of each genome in the enriched sample, after filtering by a p-value cutoff and summing over the three data partitions. The solid lines shows the resulting average sensitivity over all 400 genomes. The variability between genomes is shown as quantiles in red.