

#### Food-borne bacteria undercover: MAG reconstruction can reveal the biases in cultured food bacterial isolates and help identifying new species and strains

Sybille Tabuteau, Stéphane Chaillou

#### ► To cite this version:

Sybille Tabuteau, Stéphane Chaillou. Food-borne bacteria undercover: MAG reconstruction can reveal the biases in cultured food bacterial isolates and help identifying new species and strains. ISME18 - 18th International Symposium on Microbial Ecology, Aug 2022, Lausanne (CH), Switzerland. . hal-04283313

#### HAL Id: hal-04283313 https://hal.inrae.fr/hal-04283313v1

Submitted on 13 Nov 2023

**HAL** is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution 4.0 International License



Égalité

Fraternité



# **Food-Borne Bacteria Undercover**

MAGs (metagenome-assembled genomes) can reveal biases in cultured food bacterial isolates and help identifying new species and strains.

## Sibylle Tabuteau<sup>1</sup> and <u>Stéphane Chaillou<sup>1</sup></u>

HYPOTHESIS

Food-borne bacteria are considered non-recalcitrant to isolation on culture medium. However, recent studies on food microbiota using 16S amplicon

The dataset is composed of ~80 food metagenomes of raw sausages<sup>A</sup> and fermented vegetables<sup>B</sup>. To verify the hypothesis, 3 Target species were studied as models:

**MODELS & DATASET** 

INRA

PARIS-SACLAY

universite

sequencing has revealed that this paradigm deserves to be re-evaluated scientifically. The most obvious bias concerns the search for subdominant species (or even strains), the most part of which pass under the detection filter of isolation practices (requirement for non-canonical growth conditions, or less competitive than dominant strains), although these taxa may represent an important functional role, or participate in certain processes in a transient manner (dynamic succession of populations).

RESULTS



#### 1. Latilactobacillus sakei

Phylogenomic trees of core genomes for the three target species. Isolates are shown in colored background whereas MAGs are shown in lighter colored background. Colors indicate main phylogenetic clades. New clades can clearly be detected for each species (yellow for *L. sakei* and purple for *B. thermosphacta*). For *D. aligida* (tree N°2), average Nucleotide index revealed that some MAGs (i.e; MAGDSP in yellow background) are representing a new candidate *Dellaglioa* species.

- **1.** Latilactobacillus sakei (60 genomes available, high diversity of strains)
- Brochothrix thermopshacta (25 genomes available, low diversity of strains)
- Dellaglioa algida (4 genomes available, hard to cultivate)

1. Assembly and binning of MAGs from metagenomic data

METHODS



## 2. Dellaglioa algida



### 3. Brochothrix thermosphacta



Taxonomic assignment and reclustering of MAGs Bin 4 Unbin Bin 2 Bin 3 Unbin Bin 1 Unbin Bin 1 fastree & R scripts

ISOLATES

MAGS

2. Refining the MAGs by clustering several bins and unbinned contigs using taxonomic assignment = better recovery of genomic islands and plasmids

3. Comparative phylogenomic analysis of core and accesory genomes between MAGs and published genomes of isolates

 $\mathbf{\mathbf{v}}$ 



coregenome

#### CONCLUSIONS

> With only few metagenomes and few target species, our results clearly demonstrate the existence of strong biases in the diversity of cultured food-borne bacterial isolates.

 $\succ$  The potential of identifying new strains, clades or even species in food metagenomes is very high.



## <sup>1</sup> MICALIS INSTITUTE, INRAE Jouy-en-Josas, AgroParisTech, Université Paris-Saclay. France stephane.chaillou@inrae.fr

#### References

A Poirier et al., Data Brief. 2020 Mar 20;30:105453. doi: 10.1016/j.dib.2020.105453.

B Leech et al., mSystems. 2020 Nov 10;5(6):e00522-20. doi: 10.1128/mSystems.00522-20.

