

Modules de formation 2022









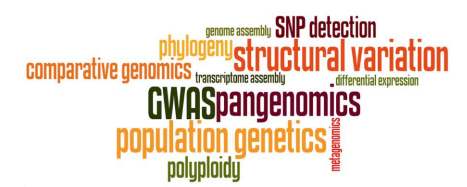








Bioinformatics platform dedicated to the genetics and genomics of tropical and Mediterranean plants and their pathogens



www.southgreen.fr









Rice

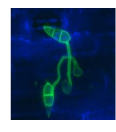
Banana

Palm









Sorghum Coffee

Cassava

Magnaporthe







Larmande Pierre

Orjuela-Bouniol Julie

Sabot François

Tando Ndomassi

Tranchant-Dubreuil Christine



Comte Aurore
Dereeper Alexis
Ravel Sébastien



Bocs Stephanie
Boizet Alice

De Lamotte Fredéric

Droc Gaetan

Dufayard Jean-François

Hamelin Chantal

Martin Guillaume

Pitollat Bertrand

Ruiz Manuel

Sarah Gautier

Summo Marilyne



Rouard Mathieu

Guignon Valentin
Catherine Breton



Sempere Guilhem











Workflow manager

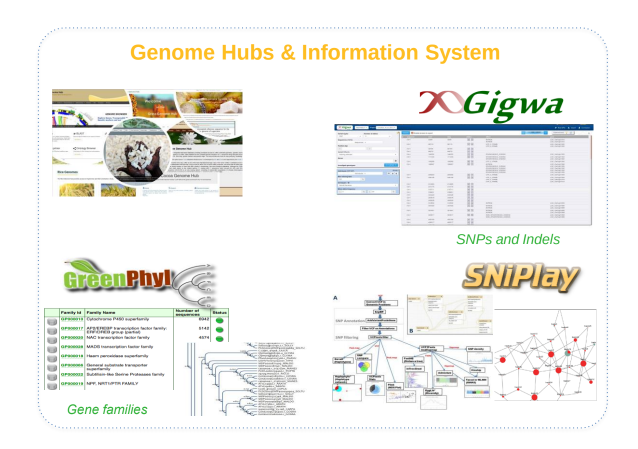






HPC and trainings....







https://github.com/SouthGreenPlatform



The South Green portal: a comprehensive resource for tropical and Mediterranean crop genomics, Current Plant Biology, 2016

I-Trop

Plant & Health Bioinformatics Platform















https://bioinfo.ird.fr/



AURORE COMTE



ALEXIS DEREEPER



BRUNO GRANOUILLAC



JULIE ORJUELA



NDOMASSI TANDO



CHRISTINE TRANCHANT

bioinfo@ird.fr



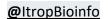
IE bioinfo































Modules de formation 2022

Toutes nos formations :

https://southgreenplatform.github.io/trainings/

Topo & TP : <u>python</u>

















Génomique Comparative Bactérienne











Two Approaches to Microbial Genomics

Starting with sets of reads representing your study isolates...



Assembly-based

- 1. Assemble each set of reads into a genome sequence
- 2. Annotate each genome
- 3. Cluster genes and compare between each genome

Variant-based

- 1. Compare each read set to a reference genome assembly
- 2. Directly compare variants between each genome

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Assembly-based

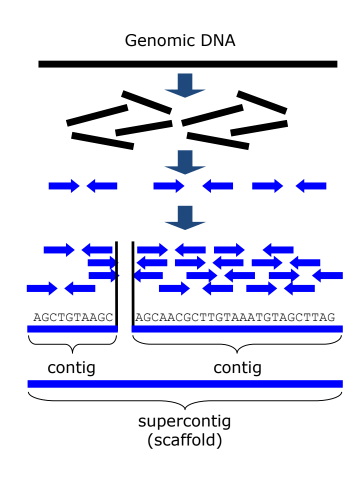
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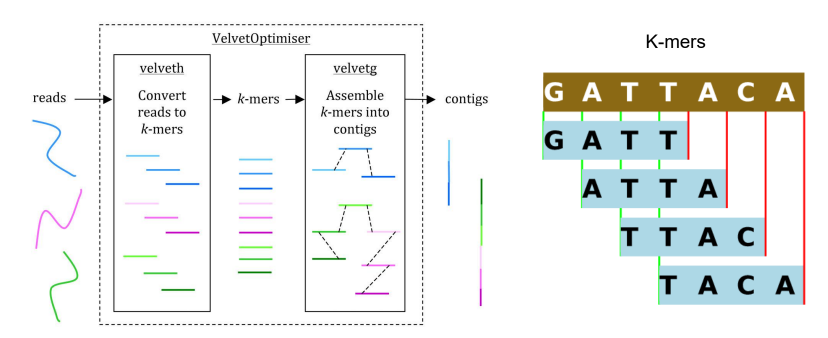
1) Assembly

Assembly Basics (de-novo assembly)



Assembly Methods

- SPAdes (http://cab.spbu.ru/software/spades/)
- Velvet (https://www.ebi.ac.uk/~zerbino/velvet/)
- Both are De Bruijn graph assemblers



Edwards and Holt 2013 MIE





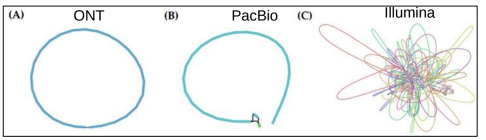
Brief Report

Comparison of De Novo Assembly Strategies for Bacterial Genomes

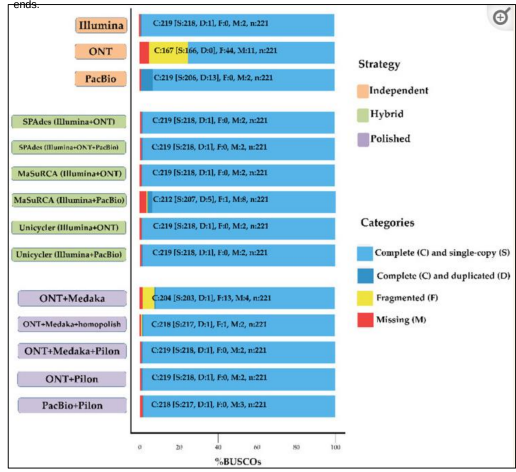
Pengfei Zhang 1,2,† Dike Jiang 1,2,†, Yin Wang 1,2,*, Xueping Yao 1,2, Yan Luo 1,2 and Zexiao Yang 1,2

Table 1					
Statistics of genome-assembly results of independent assembly strategies. $ \\$					
Platforms	Platforms Assembler Contigs Largest Contig (Largest Contig (bp)	N50	GC%
Illumina	SPAdes	527	157,573	40,498	39.87
PacBio	Canu	25	2,351,556	2,351,556	40.01
ONT	Canu	1	2,360,091	2,360,091	40.02

Table 2						
Statistics of genome-assembly results of hybrid assembly strategies.						
Assembler	Contigs	Total Length (bp)	N50	GC%		
SPAdes	266	2,402,219	1,953,224	39.97		
SPAdes	236	2,410,042	2,351,543	40.02		
Unicycler	1	2,349,186	2,349,186	40.03		
Unicycler	1	2,349,340	2,349,340	40.03		
MaSuRCA	1	2,365,339	2,365,339	40.02		
MaSuRCA	4	2,395,409	1,345,876	40.04		
	Assembler SPAdes SPAdes Unicycler Unicycler MaSuRCA	Assembler Contigs SPAdes 266 SPAdes 236 Unicycler 1 Unicycler 1 MaSuRCA 1	Assembler Contigs Total Length (bp) SPAdes 266 2,402,219 SPAdes 236 2,410,042 Unicycler 1 2,349,186 Unicycler 1 2,349,340 MaSurca 1 2,365,339	Assembler Contigs Total Length (bp) N50 SPAdes 266 2,402,219 1,953,224 SPAdes 236 2,410,042 2,351,543 Unicycler 1 2,349,186 2,349,186 Unicycler 1 2,349,340 2,349,340 MaSurca 1 2,365,339 2,365,339		

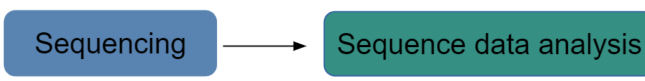


Comparison of results of independent assembly strategies. (A) Genome assembled with nanopore reads; (B) longest contig assembled with PacBio reads; (C) genome assembled with Illumina reads. Plots were obtained by using Bandage on the "assembly_graph.gfa" output file from SPAdes or the "contig.gfa" output file from Canu. Connections between contigs represent overlaps between contig



Evaluation of completeness of assembly results of different strategies. Assessments of the completeness of the assembly genomes with the datasets of proteobacteria_odb9 lineage. Bar charts produced with BUSCO plotting tool to show proportions that were classified as complete (C, blue), complete single copy (S, light blue), complete duplicated (D, dark blue), fragmented (F, yellow), and

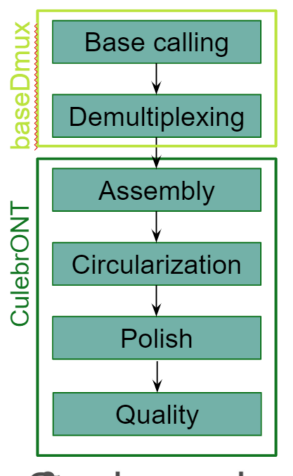
Bioinformatic Workflows: assembly











Snakemake



https://github.com/vibaotram/baseDmux



https://culebront-pipeline.readthedocs.io/en/latest/



2) Separate chromosomal and plasmid scaffolds/contigs

MICROBIAL GENOMICS

METHODS PAPER

Robertson and Nash, Microbial Genomics 2018;4 DOI 10.1099/mgen.0.000206





MOB-suite: software tools for clustering, reconstruction and typing of plasmids from draft assemblies

James Robertson¹ and John H. E. Nash^{2,*}

MOB-suite: Software tools for clustering, reconstruction and typing of plasmids from draft assemblies

Introduction

Plasmids are mobile genetic elements (MGEs), which allow for rapid evolution and adaption of bacteria to new niches through horizontal transmission of novel traits to different genetic backgrounds. The MOB-suite is designed to be a modular set of tools for the typing and reconstruction of plasmid sequences from WGS assemblies.

The MOB-suite depends on a series of databases which are too large to be hosted in git-hub. They can be downloaded or updated by running mob_init or if running any of the tools for the first time, the databases will download and initialize automatically if you do not specify an alternate database location. However, they are quite large so the first run will take a long time depending on your connection and speed of your computer. Databases can be manually downloaded from here.

Our new automatic chromosome depletion feature in MOB-recon can be based on any collection of closed chromosome sequences.

Citations

Below are the manuscripts describing the algorithmic approaches used in the MOB-suite.

- Robertson, James, and John H E Nash. "MOB-suite: software tools for clustering, reconstruction and typing of plasmids from draft assemblies." Microbial genomics vol. 4,8 (2018): e000206. doi:10.1099/mgen.0.000206
- Robertson, James et al. "Universal whole-sequence-based plasmid typing and its utility to prediction of host range and epidemiological surveillance." Microbial genomics vol. 6,10 (2020): mgen000435. doi:10.1099/mgen.0.000435

MOB-init

On first run of MOB-typer or MOB-recon, MOB-init (invoked by mob_init command) should run to download the databases from figshare, sketch the databases and setup the blast databases. However, it can be run manually if the databases need to be re-initialized OR if you want to initialize the databases in an alternative directory.

MOB-cluster

This tool creates plasmid similarity groups using fast genomic distance estimation using Mash. Plasmids are grouped into clusters using complete-linkage clustering and the cluster code accessions provided by the tool provide an approximation of operational taxonomic units OTU's. The plasmid nomenclature is designed to group highly similar plasmids together which are unlikely to have multiple representatives within a single cell and have a strong concordance with replicon and relaxase typing but is universally applicable since it uses the complete sequence of the plasmid itself rather than specific biomarkers.

MOB-recon

This tool reconstructs individual plasmid sequences from draft genome assemblies using the clustered plasmid reference databases provided by MOB-cluster. It will also automatically provide the full typing information provided by MOB-typer. It optionally can use a chromosome depletion strategy based on closed genomes or user supplied filter of sequences to ignore.

MOB-typer

Provides in silico predictions of the replicon family, relaxase type, mate-pair formation type and predicted transferability of the plasmid. Using a combination of biomarkers and MOB-cluster codes, it will also provide an observed host-range of your plasmid based on its replicon, relaxase and cluster assignment. This is combined with information mined from the literature to provide a prediction of the taxonomic rank at which the plasmid is likely to be stably maintained but it does not provide source attribution predictions.

3) Genome Annotation

Annotation Methods

- Annotation refers to assign function to DNA sequences
- There are different annotation algorithms for proteincoding genes, tRNAs, rRNAs, other non-coding RNAs
- Prokka
 (http://www.vicbioinformatics.com/software.prokka.sh tml) is an all-in-one wrapper for these tools

Table 1. Feature prediction tools used by Prokka

Tool (reference)	Features predicted
Prodigal (Hyatt 2010) RNAmmer (Lagesen <i>et al.</i> , 2007) Aragorn (Laslett and Canback, 2004) SignalP (Petersen <i>et al.</i> , 2011) Infernal (Kolbe and Eddy, 2011)	Coding sequence (CDS) Ribosomal RNA genes (rRNA) Transfer RNA genes Signal leader peptides Non-coding RNA

Then: annotate

Adding biological info to sequences

ribosome binding site

delta toxin

PubMed: 15353161

transfer RNA

Leu-(UUR)

tandem repeat

homopolymer 10 x T

Annotation

What's in an annotation?

```
    Location
```

```
• which sequence? chromosome 2
```

o where on the sequence? 100..659

what strand? -ve

Feature type

o what is it?

protein coding gene

Attributes

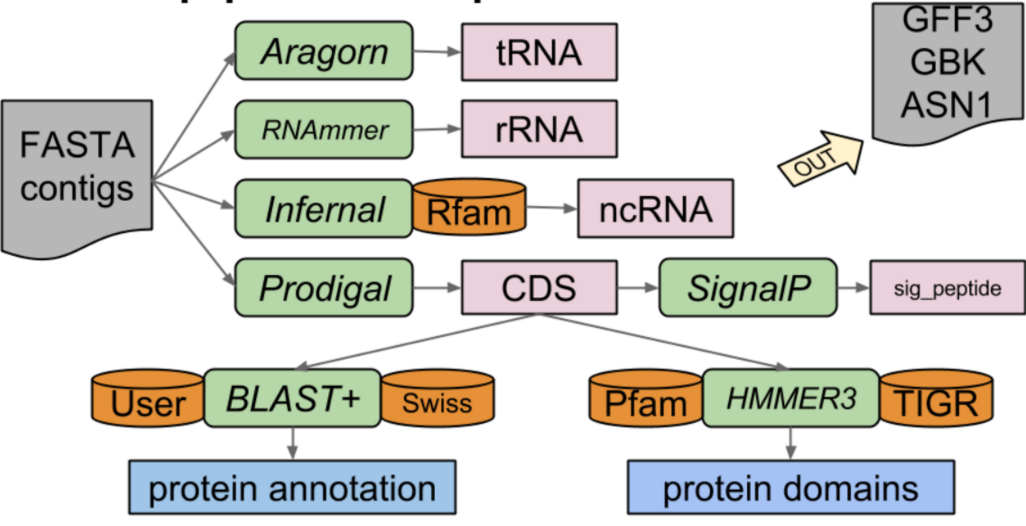
```
o protein product? alcohol dehydrogenase
```

enzyme code? *EC:1.1.1.1*

subcellular location? cytoplasm

o **note?** beer processing

Prokka pipeline (simplified)

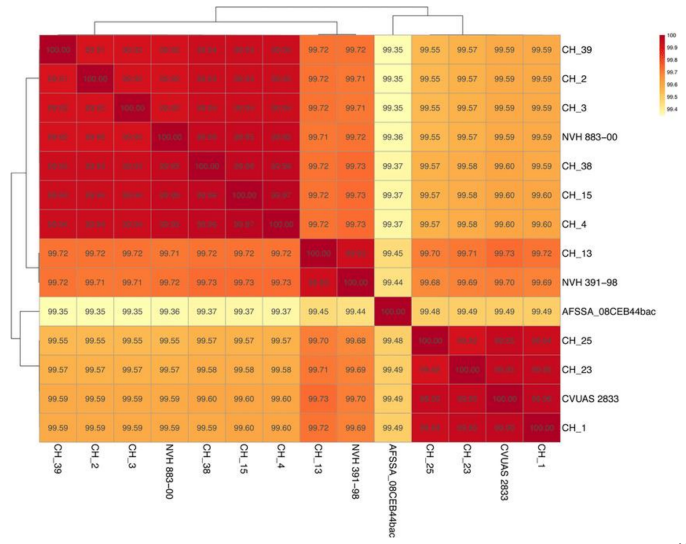


4) Pairwise Average Nucleotide Identity (ANI)

ANI: Average Nucleotide Identity

The average nucleotide identity (ANI) is a similarity index between a given pair of genomes that can be applicable to prokaryotic organisms independently of their G+C content, and a cutoff score of >95% indicates that they belong to the same species

Program: FastANI



Heat map of the average nucleotide identity (ANI) for strains of the species B. cytotoxicus (Stevens et al., 20.19)

5) Pan-genome and Gene clustering

Gene Clustering - how it works

- Assess the similarity of every gene to every other gene
 - e.g., using BLAST
- Use that similarity to join pairs of genes
 - e.g., using Reciprocal Best Hits
- Connect the gene pairs into larger clusters
 - e.g., using Reciprocal Best Hits or Markov clustering
 - => Programs: OrthoMCL, Roary, PGAP...

Table 1. Popular software for evolutionary pangenomics

Name	Authors	Reference
Panseq	Laing et al. (2010)	[12]
PanCGHweb	Bayjanov et al. (2010)	[13]
CAMBer	Wozniak et al. (2011)	[14]
PGAT	Brittnacher et al. (2011)	[15]
PGAP	Zhao et al. (2012)	[16]
GET_HOMOLOGUES	Contreras-Moreira and Vinuesa (2013)	[17]
GET_HOMOLOGUES-EST	Contreras-Moreira et al. (2017)	[18]
PanTools	Sheikhizadeh et al. (2016)	[19]
EDGAR 2.0	Blom et al. (2016)	[20]
PanX	Ding et al. (2018)	[21]
Micropan	Snipen and Liland (2015)	[22]
FindMyFriends	Pedersen (2015)	[23]
Piggy	Thorpe et al. (2018)	[24]
PanViz	Pedersen et al. (2017)	[25]

Method	Software	Input	Graph output	Pan-genome	Sequence homology	Paralogue identification
Roary (v3.13.0)	Conda package	GFF3	DOT	Directed graph	BLAST	Synteny
Ptolemy (v1.0)	Java executable	FASTA+GFF	GFA	Directed graph	minimap2	Graph-based
PPanGGoLin (v1.0.13)	Conda package	GBK or FASTA	GEXF	Undirected graph	MMseq2	Synteny
PIRATE (v1.0.3)	Conda package	GFF3	GFA	Directed graph	BLAST (/DIAMOND)	Synteny
Panaroo (v1.1.2)	Conda package	GFF3	GML	Directed graph	CD-HIT	Synteny

MICROBIAL GENOMICS

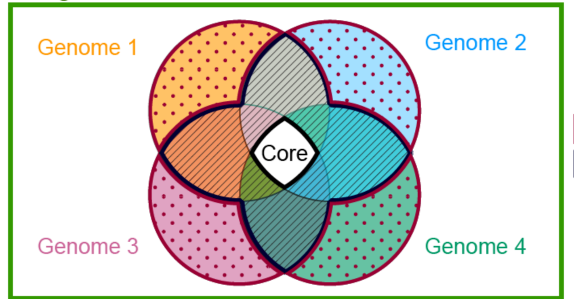
Volume 7, Issue 11

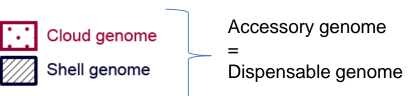
Research Article | Open Access

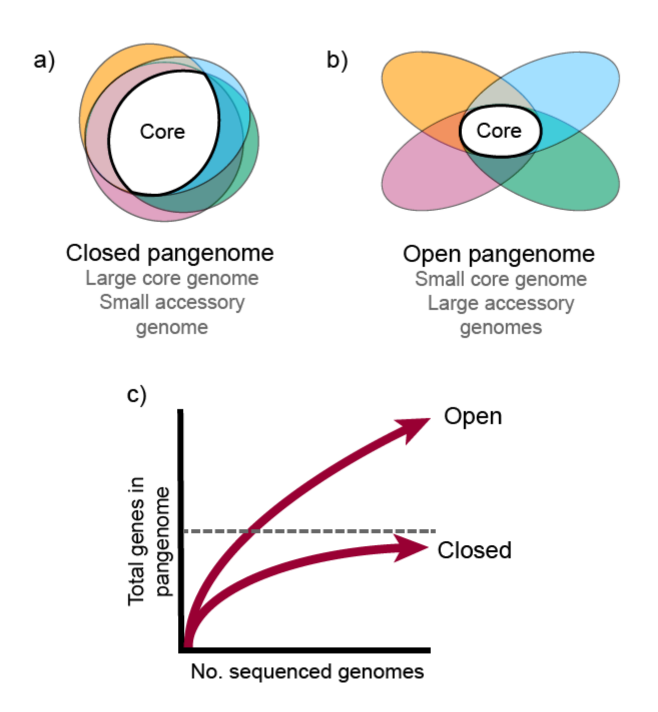
A comparative study of pan-genome methods for microbial organisms: Acinetobacter baumannii pan-genome reveals structural variation in antimicrobial resistance-carrying plasmids 3

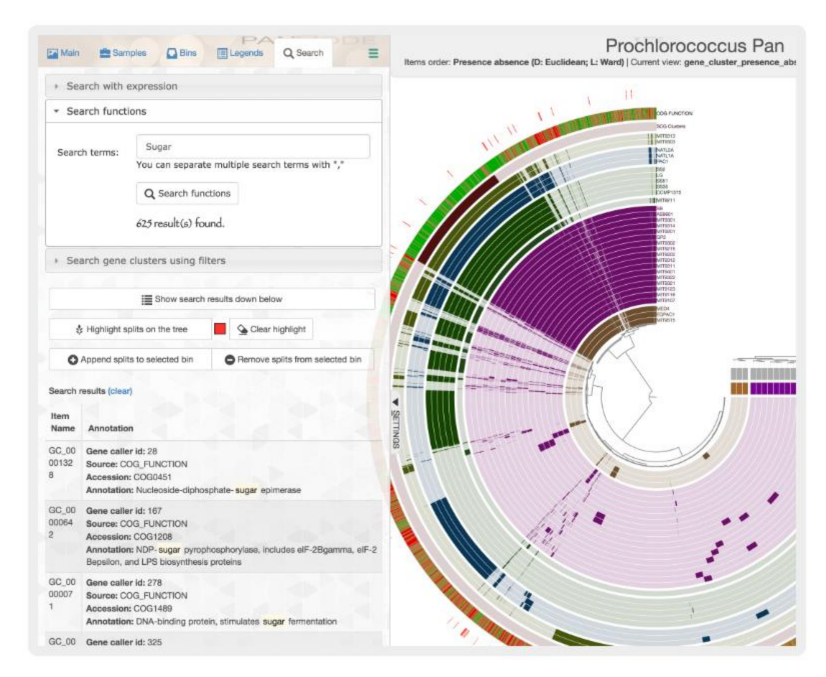
Aysun Urhan¹, Thomas Abeel^{1,2}

Pangenome

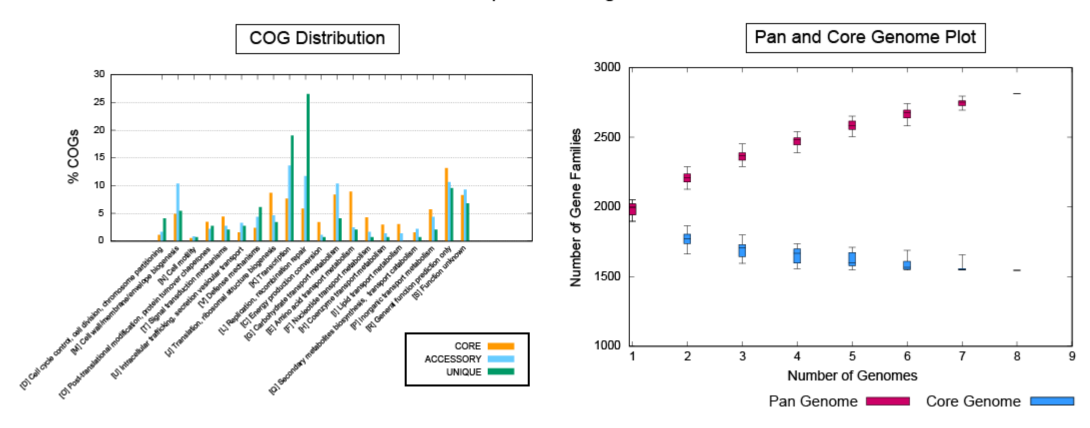








BPGA (Bacterial Pan Genome Analysis tool) Streptococcus agalactiae



6) Pan-GWAS

Pan-GWAS

Pan-GWAS of *Streptococcus agalactiae* Highlights Lineage-Specific Genes Associated with Virulence and Niche Adaptation

Authors: Andrea Gori , Odile B. Harrison, Ethwako Mlia, Yo Nishihara, Jia Mun Chan, Jacquline Msefula, Macpherson Mallewa, SHOW ALL (13 AUTHORS), Robert S. Heyderman | AUTHORS INFO & AFFILIATIONS

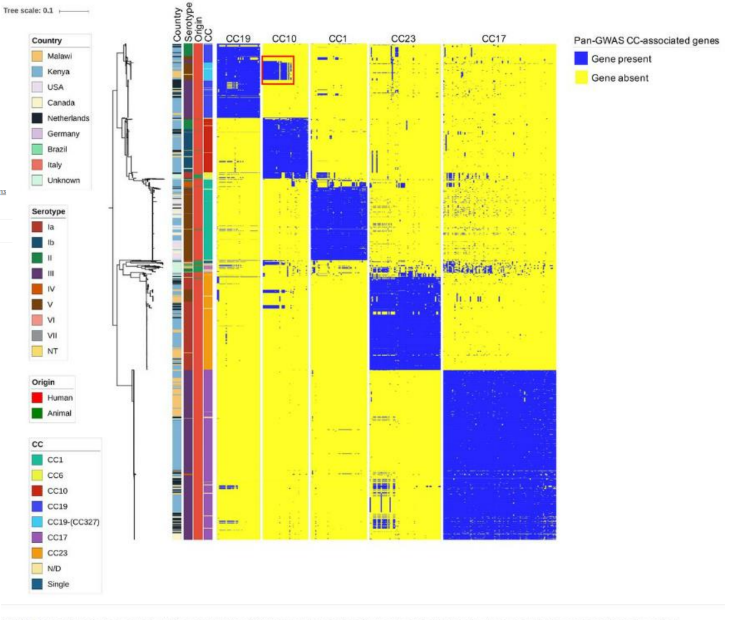


FIG 2 Core genome-based population structure of GBS. The phylogenetic tree is annotated with 4 colored strips representing the clonal complex, the country of isolation, the origin, and the serotype of each strain. The three binary heatmaps represent the presence (blue) or absence (yellow) of the genes identified by the pan-GWAS pipeline. The tree is rooted at midpoint. The reference strain used in this analysis was COH1, reference HC939456. The red square in the CC10 heatmap highlights the cluster of CC10-associated genes found in CC19 clones. Trees built with different reference strains are shown in Fig. S1 in the supplemental material and show analogous topology.

Un odds ratio:

- < 1 signifie que l'événement est moins fréquent dans le groupe A que dans le groupe B;
- = 1 signifie que l'événement est aussi fréquent dans les deux groupes ;
- > 1 signifie que l'événement est plus fréquent dans le groupe A que dans le groupe B.



Merci pour votre attention!



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SUIVEZ NOUS SUR TWITTER!





South Green: ogreen_bioinfo



I-Trop: <u>@ItropBioinfo</u>



N'oubliez pas de nous citer!

Comment citer les clusters?

"The authors acknowledge the IRD i-Trop HPC at IRD Montpellier for providing HPC resources that have contributed to the research results reported within this paper. URL: http://bioinfo.ird.fr/ "

"The authors acknowledge the CIRAD UMR-AGAP HPC (South Green Platform) at CIRAD montpellier for providing HPC resources that have contributed to the research results reported within this paper. URL:

http://www.southgreen.fr"