

## Training on Galaxy: Metabarcoding October 2024 - Webinar

## **FROGS Practice on function inference**

**Gigenae** GenPhySE 🗫 MaiAGE GAB

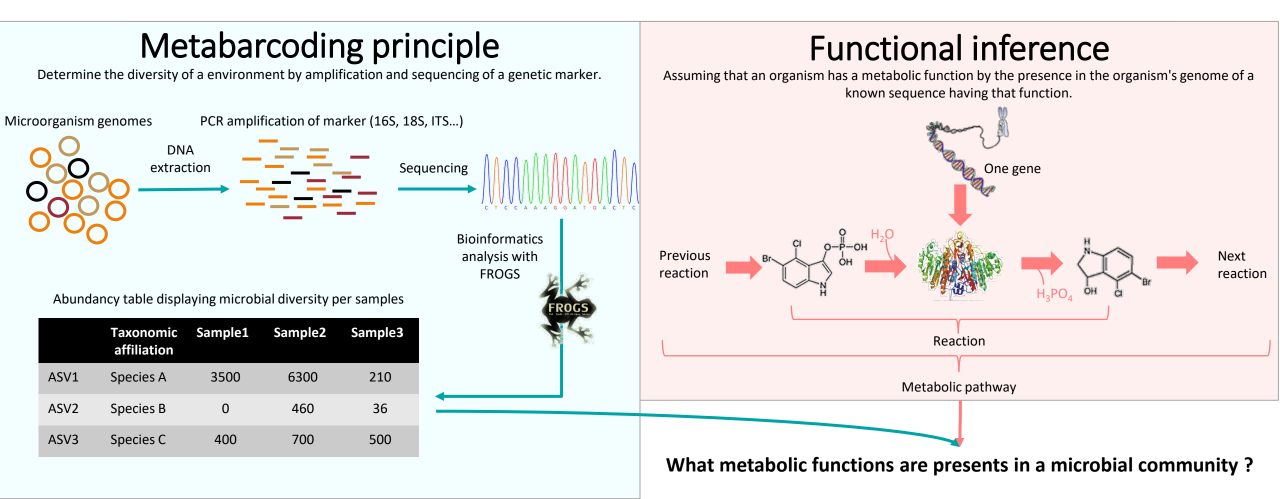
LUCAS AUER, MARIA BERNARD, LAURENT CAUQUIL, VINCENT DARBOT, MAHENDRA MARIADASSOU, GÉRALDINE PASCAL & OLIVIER Rué

micipile



# What metabolic functions are present in the environment?

Concepts



### Based on PICRUSt2

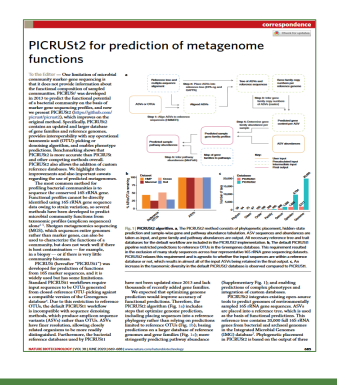
PICRUSt (Phylogenetic investigation of communities by reconstruction of unobserved states) is an open-source tool.

It is a software for predicting functional abundances based only on marker gene sequences.

PICRUSt2 is composed of 4 python applications.

No graphic interface exists to run PICRUSt2 for non-expert users.

Douglas, G.M., Maffei, V.J., Zaneveld, J.R. *et al.* **PICRUSt2** for prediction of metagenome functions. *Nat Biotechnol* 38, 685–688 (2020). <u>https://doi.org/10.1038/s41587-020-0548-6</u>

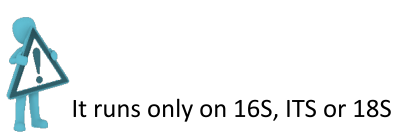


How it works ?

- 1. Places the ASVs into a reference phylogenetic tree and predicts of marker copy number in each ASV.
- 2. Predicts number of function copy number in each ASV and calculates functions abundances in each sample and ASV abundances according to marker copy number.
- 3. Calculates pathway abundances in each sample.

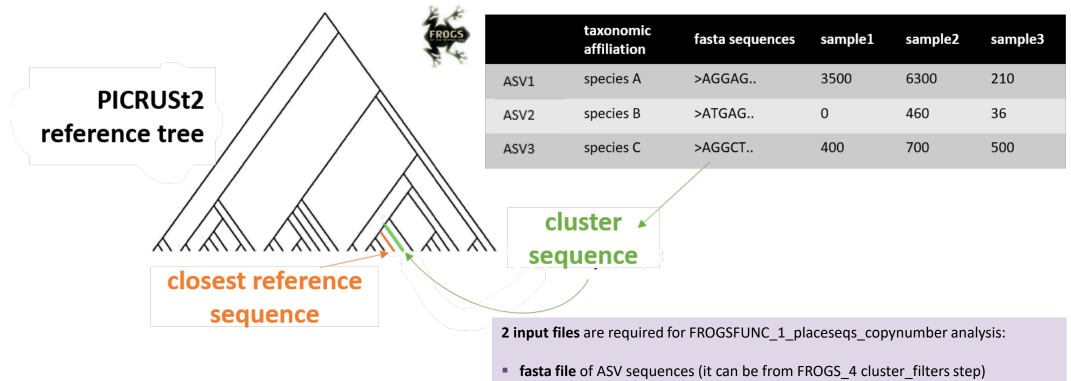
FROGSFUNC\_1\_placeseqs\_copynumber

FROGSFUNC\_2\_functions

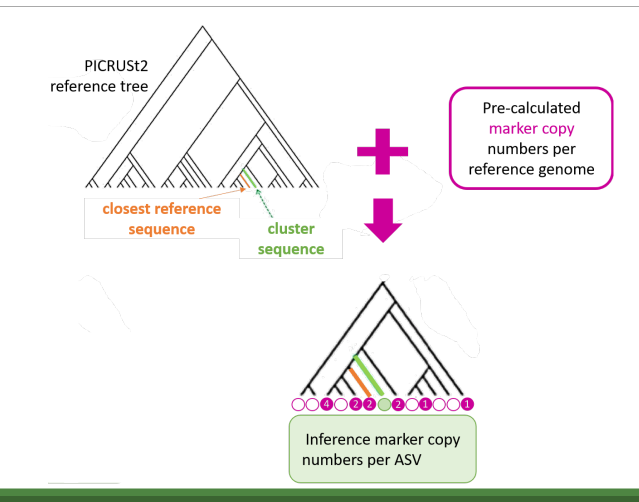


FROGSFUNC\_3\_pathways

- **FROGSFUNC\_1\_placeseqs\_copynumber** is the first step of PICRUSt2.
- It inserts your studied sequences into a <u>reference tree</u>.
- By default, this reference tree is based on 20,000 16S sequences from genomes in the Integrated Microbial Genomes database.
- Prediction of the copy numbers of the marker gene (16S, ITS or 18S) in order to normalize the ASV abundances table thereafter.



 biom file of ASV abundances with taxonomic affiliation information (it can be from FROGS\_5\_taxonomic\_affiliation step)



FR	OGSFU	JNC_1_	placeseqs_and_copynu	mbers Places ASVs	into a reference phyloger	netic tree. (Galaxy Version 4.1.0+galaxy1)	☆ Favorite	& Versions	- Optio	ns	
Seque	ence fi	le									
٥	¢		15: FROGS_4 Cluster f	ilters: clusterFilters_	sequences.fasta				•	⊳	
The se		e file to	analyse (format: fasta).	(input-fasta)							
0	ſĴ		25: FROGS_5 Taxonon	nic affiliation: affilia	tion_abundance.biom				•		
	rs 8S omic n		of interest. (ref-dir)								
	ment t	ool									
⊘ e O se	pa-ng epp					epa-ng is very memory	and com	puting po <sup>,</sup>	wer int	tensive	5
Placer (plac			insertion of sequences in	ito the reference tre	e. SEPP is a low-memory	alternative to EPA-ng for placing sequenc	es, and is only a	vailable for 16S a	nalysis.		
Minim	num ali	ignmen	nt length								
0.8	-		I,	gnore this p	arameter, it wi	ll disappear in FROGS v 5	.0	-43			
Propo <b>Email</b>			tal length of an input se	quence that must a	lign with reference seque	nces. All others will be out. (default: 0.80)	(min-align)	1			
	No										

Send an email notification when the job completes.

✓ Execute

epa-ng is the only choice

Since epa-ng is greedy then it may force to reduce the dataset to

for ITS and 18S

run.

## Input files

- Sequences file: The ASV fasta sequence file.
- biom file: The ASV <u>biom</u> file. Taxonomic affiliations must be done before (biom file form FROGS\_5\_taxonomic\_affiliation tool).
- **taxonomy marker**: 16S, ITS and 18S only available.



If your ASVs are based on another marker, you cannot use this tool.

- placement tool: EPA-NG or SEPP are placement tools for insertion of sequences into the PICRUSt2 reference tree. SEPP is a low-memory alternative to EPA-ng for placing sequences. So, if the tool crashes with EPA-ng, try again with SEPP.
- minimum alignment length: Proportion of the total length of an input sequence that must align with reference sequences. All other will be out.

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_marker.tsv

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs.biom

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_closests\_ref\_sequences.txt

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs.fasta

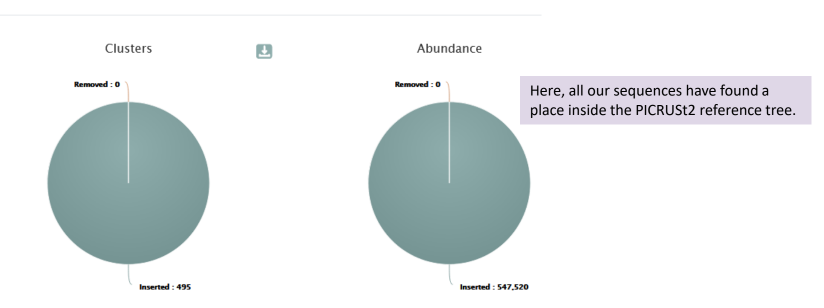
FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_excluded.tsv

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_tree.nwk

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: report.html

### FROGSFUNC\_1: report.html

#### Insertion in reference tree summary



The html report file describes that ASVs are contained or not in the phylogenetic tree.

Note that PICRUSt2 uses its own reference tree to affiliate ASVs from reference sequences.

The report file indicates for each ASV which is the closest PICRUSt2 reference sequence, and compares it to the original FROGS taxonomy. Clicking on the sequence ID gives you more information about it <u>JGI</u> database.

ASV ↑↓	Nb sequences †↓	FROGS Taxonomy	PICRUSt2 closest ID (JGI) 14	PICRUStZ closest reference name 14
luster_1	84849	Bacteria; Firmicutes; Bacilli; Lactobacillales; Listeriaceae; Brochothrix; Brochothrix thermosphacta	2576861686	Brochothrix thermosphacta FSL F6-1036
Cluster_10	4188	${\sf Bacteria}; {\sf Bacteroidota}; {\sf Bacteroidia}; {\sf Flavobacteriales}; {\sf Flavobacteriaceae}; {\sf Flavobacterium}; {\sf Flavobacterium} \ {\sf sp.}$	2724679776	Flavobacterium sp. 9
Cluster_100	696	Bacteria; Proteobacteria; Gamma proteobacteria; Xan thomonadales; Xan thomonadaceae; Stenotrophomonas; Stenotrophomonas sp.	2639762796	Stenotrophomonas maltophilia OC194
Cluster_101	752	${\sf Bacteria}; {\sf Bacteroidota}; {\sf Bacteroidia}; {\sf Flavobacteriales}; {\sf Weeksellaceae}; {\sf Chryseobacterium}; {\sf Chryseobacterium} sp.$	2600255101	Chryseobacterium haifense DSM 19056

inserted	l in the reference t	sest reference sequence from the ASV cree under the following format: der;Family;Genus;Species				Lowest same taxonomic rank		
	PICRUSt2 closest reference name 11	♥ PICRUSt2 closest taxonomy	†↓	NSTI 岸	NSTI Confidence ↑↓	between FROGS and PICRUSt2 ↑↓	Comment †↓	
	Brochothrix thermosphacta FSL F6-1036	Bacteria; Firmicutes; Bacilli; Bacillales; Listeriaceae; Brocho	othrix;Brochothrix thermosphacta	0.062	Good	Up to Species	/	
	Flavobacterium sp. 9	Bacteria; Bacteroi detes; Flavobacteriia; Flavobacteriales;	Flavobacteriaceae;Flavobacterium;Flavobacterium sp.	0.057	Good	Up to Species	/	
	Stenotrophomonas maltophilia OC194	Bacteria; Proteobacteria; Gammaproteobacteria; Xantho maltophilia	monadales;Xanthomonadaceae;Stenotrophomonas;Stenotrophomonas	Nearest Sequenced Taxon Index ( <u>NSTI</u> ) is the phylogene distance between the ASV and the nearest sequenc reference genon				
	Chryseobacterium haifense DSM 19056	Bacteria; Bacteroi detes; Flavobacteriia; Flavobacteriales;	Weeksellaceae;Kaistella;Chryseobacterium haifense	0.063	Good	Up to Family	/	

PICRUSt2 closest reference name 14	PICRUSt2 closest taxonomy 14	NSTI †↓	NSTI Confidence †4	Lowest same taxonomic rank between FROGS and PICRUSt2 14	Comment ↑↓
Brochothrix thermosphacta FSL F6-1036	Bacteria; Firmicutes; Bacilli; Bacillales; Listeriaceae; Brochothrix; Brochothrix thermosphacta	0.062	Good	Up to Species	/
Flavobacterium sp. 9	Bacteria; Bacteroidetes; Flavobacteriia; Flavobacteriales; Flavobacteriaceae; Flavobacterium; Flavobacterium sp.	0.057	Good	Up to Species	/
Stenotrophomonas maltophilia OC194	Bacteria; Proteobacteria; Gammaproteobacteria; Xanthomonadales; Xanthomonadaceae; Stenotrophomonas; Stenotrophomonas maltophilia	0.036	Good	Up to Genus	/
Chryseobacterium haifense DSM 19056	Bacteria; Bacteroidetes; Flavobacteriia; Flavobacteriales; Weeksellaceae; Kaistella; Chryseobacterium haifense	0.063	Good	Up to Family	/



According to the NSTI score, we guide you in the confidence you can bring to the issue affiliation of PICRUSt2. Four levels are given:

- 0 < Good < 0.5
- 0.5 <= Medium < 1
- $1 \le Bad \le 2$
- To exclude  $\geq 2$

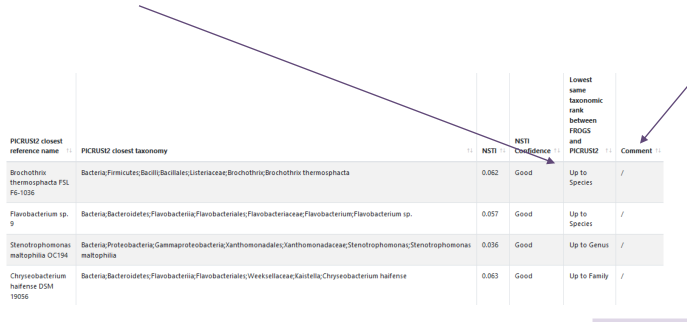
PICRUSt2 sets NSTI threshold to 2 per default. Some studies have shown that this threshold is permissive. Thus, it is important to see if the taxonomies between PICRUSt2 and FROGS are quite similar or not, in order to potentially choose a more stringent threshold afterwards.

For example, a NSTI lower than 0.5, with "species" as lowest common taxonomic rank between FROGS and PICRUSt2 will product a good prediction.

PICRUSt2 closest reference name 14	PICRUSt2 closest taxonomy 14	NSTI †4	NSTI Confidence ↑↓	Lowest same taxonomic rank between FROGS and PICRUSt2 14	Comment †↓
Brochothrix thermosphacta FSL F6-1036	Bacteria; Firmicutes; Bacilli; Bacillales; Listeriaceae; Brochothrix; Brochothrix thermosphacta	0.062	Good	Up to Species	/
Flavobacterium sp. 9	Bacteria; Bacteroidetes; Flavobacteriia; Flavobacteriales; Flavobacteriaceae; Flavobacterium; Flavobacterium sp.	0.057	Good	Up to Species	/
Stenotrophomonas maltophilia OC194	Bacteria; Proteobacteria; Gammaproteobacteria; Xanthomonadales; Xanthomonadaceae; Stenotrophomonas; Stenotrophomonas maltophilia	0.036	Good	Up to Genus	/
Chryseobacterium haifense DSM 19056	Bacteria; Bacteroidetes; Flavobacteriia; Flavobacteriales; Weeksellaceae; Kaistella; Chryseobacterium haifensen en senten en	0.063	Good	Up to Family	/

#### Lowest same taxonomic rank between FROGS and PICRUSt2 : Lowest

common taxonomic rank between FROGS and PICRUSt2 affiliations.



#### Comment :

- *identical taxonomy*: if the FROGS and
   PICRUSt2 taxonomic affiliations are identical.
- *identical sequence*: if the ASV sequence is strictly the same as the reference sequence.
- it is a mark of unambiguity



Search « Up to Species » for obtaining less ambigous reference

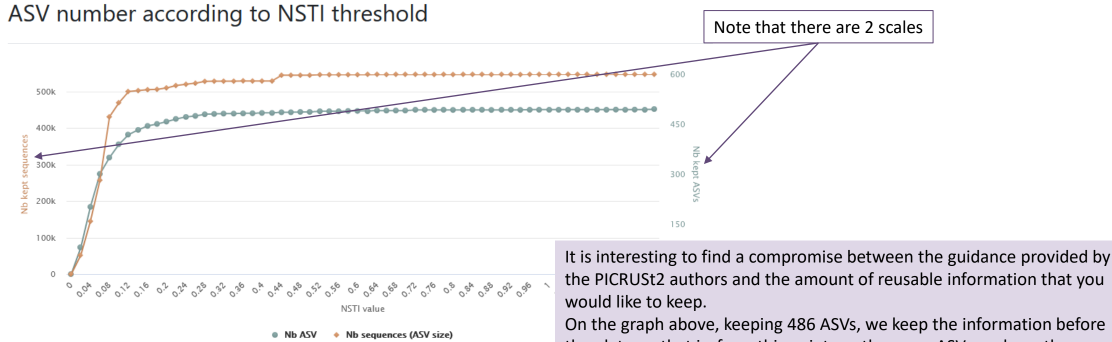
PICRUSt2 closest	PICRUSt2 closest taxonomy	NSTI †4	NSTI Confidence †1	Lowest same taxonomic rank between FROGS and PICRUSt2 14	Comment 14
Brochothrix thermosphacta FSL F6-1036	${\sf Bacteria}; {\sf Firmicutes}; {\sf Bacillaes}; {\sf Listeriaceae}; {\sf Brochothrix}; {\sf Brochothrix} thermosphactable is {\sf Bacteria}; {\sf Bacillaes}; {\sf Listeriaceae}; {\sf Brochothrix}; {\sf Brochothrix}; {\sf Brochothrix}; {\sf Brochothrix}; {\sf Bacteria}; {\sf Bacillaes}; {\sf Brochothrix}; {\sf Brochothrix}; {\sf Brochothrix}; {\sf Bacillaes}; {\sf Bacil$	0.062	Good	Up to Species	1
Flavobacterium sp. 9	${\sf Bacteria}; {\sf Bacteroidetes}; {\sf Flavobacteria}; {\sf Flavobacteria}; {\sf Flavobacteria}; {\sf Flavobacterium}; {\sf Flavobacter$	0.057	Good	Up to Species	1
Stenotrophomonas maltophilia OC194	Bacteria; Proteo bacteria; Gamma proteo bacteria; Xanthomona dales; Xanthomona daceae; Stenotrophomonas; Stenotrophomonas maltophilia	0.036	Good	Up to Genus	/
Chryseobacterium haifense DSM 19056	Bacteria;Bacteroidetes;Flavobacteriia;Flavobacteriales;Weeksellaceae;Kaistella;Chryseobacterium haifense	0.063	Good	Up to Family	/

PICRUSt2 reference tree is base on NCBI taxonomy. If you want more « Up to Sepices », i.e. more correspondances between FROGS affiliation and PICRUSt2 affiliation , think to use 16S REFseq databank in FROGS\_5 Taxonomic affiliation step

FROGS_5 Taxonomic affiliation 1	xonomic affiliation of each ASV's seed by RDPtools and BL
Using reference database	-``@`-
16S REFseq Bacteria 20230726	
Select reference from the list	

For this exemple, affiliation with 16S SILVA 138.1 gives 73 « Up to Species » and with 16S REFseq 20230726 gives 146 « Up to Species »

### Another key for choosing NSTI threshold



N.B.: Select area to zoom in.

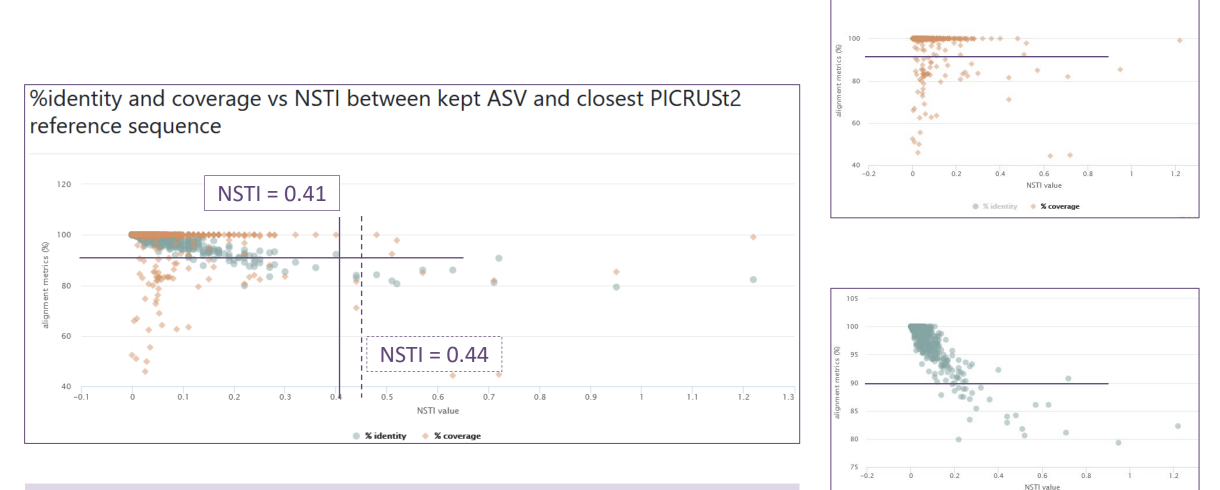
the PICRUSt2 authors and the amount of reusable information that you

On the graph above, keeping 486 ASVs, we keep the information before the plateau, that is, from this point on, the more ASVs we keep the more we degrade the accuracy. So, here NSTI = 0.44 But this depends strongly on the datasets.



This graph allows you to set the "NSTI cut-off" parameter of the next tool

### Another key for choosing NSTI threshold



120

With %id and %cov cutoff at 90%, we can choose a NSTI cutoff at 0.41 or 0.44, we will select same data. So we can choose 0.41.

% identity

♦ % coverage

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_marker.tsv

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs.biom

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_closests\_ref\_sequences.txt

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs.fasta

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_excluded.tsv

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_tree.nwk

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: report.html

### FROGSFUNC\_1: tree.nwk

(2609460310:0.0708,2713896746:0.079553):0.020861):0.018755):0.049721,2667528167:0.021242):0.05 (2634166173:0.020087,((2630968917:0.042097,(2695420983:0.0414,((641380439:0.006668,(275118574¢ (2630968293:0.07529,(2634166307:0.010503,2545824761:0.03788):0.019526):0.024717):0.015629):0.¢ ((2630968881:0.054077,2654587584:0.013434):0.046038,2585427602:0.014665):0.024954):0.022354):¢ ((2526164557:0.215838,((2524023070:0.051755,(2627853707:0.116778,2509276063:0.03521):0.033363) ((2627853601:0.074019,

((640963037:0.107024,2675903215:0.024681):0.038088,2728369219:0.027685):0.020008):0.023448,274
0.021443):0.036439,((((2600255390:0.00005,Cluster\_103:0.022437):0.088344,(2617271337:0.049512,
(2585427837:0.000001,2693429891:0.001563):0.000001):0.00005,Cluster\_29:0.031725):0.013777):0.@
(((((2700988729:0.065717,2695420957:0.031748):0.022352,(2698536752:0.009476,(((((2695421021:0.
((2643221790:0.0157,2728368985:0.008527):0.014802,(2695420960:0.010196,Cluster\_188:0.040768):@
(2738541279-cluster:0.000001,((2636415472:0.00924,(2648501293:0.00005,Cluster\_202:0.029009):0.
((((((2734482184:0.025815,2648501732:0.010421):0.005141,

(2513020052:0.021475,2695420959:0.057031):0.010852):0.005258,2754412712:0.020212):0.011771,265
((2700988712:0.032914,2695420925-cluster:0.003371):0.007089,(((2619618807:0.048997,(2523533607)
(2693429909:0.000001,Cluster\_192:0.14387):0):0.017851,((2684622654:0.032648,(2551306066:0.0026)
(2693429883:0.000001,Cluster\_321:0.05075):0):0.019173):0.006793):0.012944):0.039508):0.029699,
(2523533550:0.002444,Cluster\_37:0.02644):0.010937):0.020635):0.038474):0.000001,2643221667:0.6
(((2739367857-cluster:0.001694.2585427836:0.026929):0.008262.((((2619618994:0.089002.((25246)))))))))

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_marker.tsv

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs.biom

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_closests\_ref\_sequences.txt

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs.fasta

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_excluded.tsv

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_tree.nwk

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: report.html

This file contains all the ASVs that could not be placed in the PICRUSt2 reference tree.

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_marker.tsv

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs.biom

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_closests\_ref\_sequences.txt

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs.fasta

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_excluded.tsv

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_tree.nwk

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: report.html

This file contains all the ASVs fasta file (without those that may be excluded)

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_marker.tsv

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs.biom

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_closests\_ref\_sequences.txt

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs.fasta

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_excluded.tsv

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_tree.nwk

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: report.html

In this table, you will find all the details of the correspondences between your sequences and those of PICRUSt2.

### FROGSFUNC\_1: closest\_ref\_sequences.txt

#Cluster	Nb sequences	FROGS Taxonomy	PICRUSt2 closest ID
Cluster_1	84849	Bacteria;Firmicutes;Bacilli;Lactobacillales;Listeriaceae;Brochothrix;Brochothrix thermosphacta	2576861686
Cluster_2	31333	Bacteria; Proteobacteria; Gamma proteobacteria; Enterobacterales; Vibrionaceae; Photobacterium; unknown species and the second structure of the seco	2724679053
Cluster_3	40711	Bacteria; Firmicutes; Bacilli; Lactobacillales; Lactobacillaceae; Latilactobacillus; Lactobacillus sakei	2728369693
Cluster_4	22275	Bacteria; Actinobacteriota; Actinobacteria; Propionibacteriales; Propionibacteriaceae; Cutibacterium; unknown species	2537562124
Cluster_5	29355	Bacteria; Firmicutes; Bacilli; Lactobacillales; Lactobacillaceae; Leuconostoc; Leuconostoc inhae KCTC 3774	641522636

PICRUSt2 closest reference name	PICRUSt2 closest taxonomy
Brochothrix thermosphacta FSL F6-1036	Bacteria;Firmicutes;Bacilli;Bacillales;Listeriaceae;Brochothrix;Brochothrix thermosphacta
Photobacterium kishitanii 201212X	Bacteria; Proteobacteria; Gamma proteobacteria; Vibrionales; Vibrionaceae; Photobacterium; Photobacterium kishitanii
Lactobacillus curvatus JCM 1096, DSM 20019	${\sf Bacteria}; {\sf Firmicutes}; {\sf Bacilli}; {\sf Lactobacillales}; {\sf Lactobacillaceae}; {\sf Latilactobacillus}; {\sf $
Cutibacterium acnes SK182	Bacteria; Actino bacteria; Actino mycetia; Propioni bacteriales; Propioni bacteria ceae; Cutibacterium; Cutibacterium acnes action activity of the second
Leuconostoc citreum KM20	Bacteria; Firmicutes; Bacilli; Lactobacillales; Lactobacillaceae; Leuconostoc; Leuconostoc citreum

NSTI	NSTI Confidence	FROGS and PICRUSt2 lowest same taxonomic rank	Comment	Cluster sequence	PICRUSt2 closest reference sequence	%id	%cov	score
0.062	Good	Up to Species	1	GACGAACGCTGGCGGCGTC	TTAACGAGAGTTTGATCCTGGCTCAGGACGAACGCTG	100.0	100.0	493.5
0.05	Good	Up to Genus	1	ATTGAACGCTGGCGGCAGG	GAGTAATGCCTGGGAATATACCCTGATGTGGGGGGATAA	99.75	81.91	400.0
0.068	Good	Up to Genus	1	GACGAACGCTGGCGGCGTC	TTTTAATCGAGAGTTTGATCCTGGCTCAGGACGAACGC	98.84	83.27	412.0
0.039	Good	Up to Genus	identical sequence	GACGAACGCTGGCGGCGTC	TTCCATTGGAGAGTTTGATCCTGGCTCAGGACGAACG	100.0	100.0	468.0
0.072	Good	Up to Genus	1	GATGAACGCTGGCGGCGTG	GAGAGTTTGATCCTGGCTCAGGATGAACGCTGGCGGC	97.17	100.0	444.5

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_marker.tsv

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs.biom

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_closests\_ref\_sequences.txt

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs.fasta

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_excluded.tsv

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_tree.nwk

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: report.html

Abundance table without those that may be excluded

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_marker.tsv

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs.biom

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_closests\_ref\_sequences.txt

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs.fasta

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_excluded.tsv

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: frogsfunc\_placeseqs\_tree.nwk

FROGSFUNC\_1\_placeseqs\_and\_copynumbers: report.html

metadata_NSTI	16S_rRNA_Count	sequence				
0.062403	1	Cluster_1				
0.057594000000000006	1	Cluster_10				
0.036314	1	Cluster_100				
0.063955	1	Cluster_101				
0.019805	1	Cluster_102	FROGSFUNC 1: marker.tsv			
0.022487	1	Cluster_103				
0.074696	1	Cluster_104				
0.0042	7	Cluster_105				
0.048352	1	Cluster_106				
0.295495	1	Cluster_107				
0.018517	1	Cluster_108				
0.017555	1	Cluster_109				
0.017435	1	Cluster_11				
0.06880800000000001	1	Cluster_110				
0.031835	1	Cluster_111				
0.11071900000000001	1	Cluster_112				
0.0097209999999999999	1	Cluster_113	Prediction by PICRUSt2 of 16S copy number for each ASV			
0.043952	4	Cluster_114	(placed in the reference tree)			
0.105228	1	Cluster_115				
0.238852	1	Cluster_116				
0.006305	1	Cluster_117				
0.035066	1	Cluster_118				
0.0531559999999999995	1	Cluster_119				
0.060149	1	Cluster_12				
0.194218	1	Cluster_120				
0.111322	1	Cluster_121				
0.031979	1	Cluster_122				
0.037387000000000004	2	Cluster_123				
0.266484999999999997	1	Cluster_124				
0.091262	2	Cluster_125				
0.051895000000000004	1	Cluster_126				
0.019323	6	Cluster_127				
0.022711000000000002	4	Cluster_128				

How it works ?

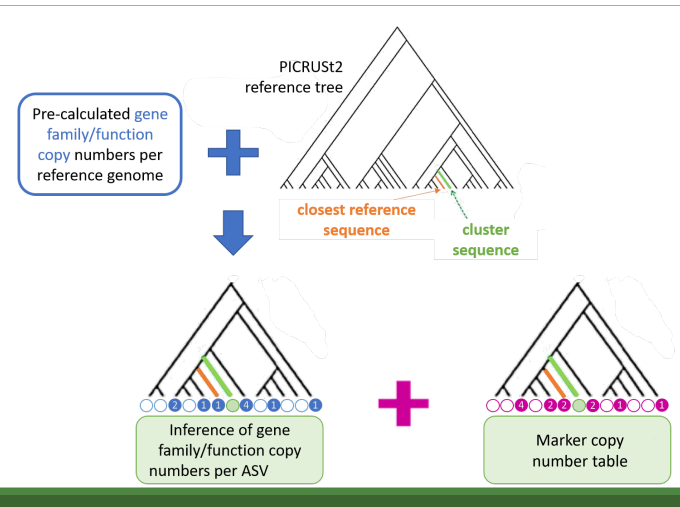
- 1. Places the ASVs into a reference phylogenetic tree and predicts of marker copy number in each ASV.
- 2. Predicts number of function copy number in each ASV and calculates functions abundances in each sample and ASV abundances according to marker copy number.
- 3. Calculates pathway abundances in each sample.

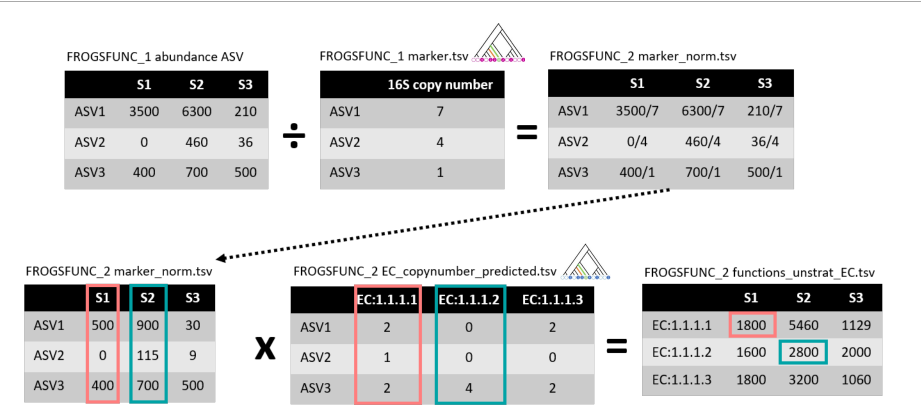
It runs only on 16S, ITS or 18S

FROGSFUNC 2 functions

FROGSFUNC\_3\_pathways







FROGSFUNC_2_functions       Calculates functions abundances in each         sample. (Galaxy Version 4.1.0+galaxy1)       Sample Versions
Biom file
Image: Construction of the second
The abundance file i.e. FROGSFUNC_1_placeseqs_copynumber tool output file (frogsfunc_placeseqs.biom). (input-biom)
Sequence file
L       L       49: FROGSFUNC_1_placeseqs_and_copynumbers: frogsfunc_placeseqs.fasta       ►
The fasta file i.e. from FROGSFUNC_1_placeseqs_copynumber tool output file (frogsfunc_placeseqs.fasta). (input-fasta)
Tree file Thanks to the previous prediction of the copy
□       □       47: FROGSFUNC_1_placeseqs_and_copynumbers: frogsfunc_placeseqs_tree.nwk       -       □       numbers of the marker gene (16S, ITS or 18S) in
The file contains the tree information from FROGSFUNC_1_placeseqs_copynumber tool (frogsfunc_placeseqs_tree.nwk). FROGSFUNC_1, FROGSFUNC_2 can normalize
the ASV abundances table.
Image: Second
Table of predicted marker copy number i.e. FROGSEUNC_1_placeseqs_copynumber output (frogsfunc_marker.tsv). (input- marker)
Taxonomic marker
⊘ 165
O ITS O 18S
Taxonomic marker of interest.

#### Prediction of the **functions abundances**, using different databases:

- EC : <u>https://enzyme.expasy.org/</u>
- KO: <u>https://www.genome.jp/kegg/ko.html</u>
- PFAM : <u>http://pfam.xfam.org/</u>
- COG : <u>https://www.ncbi.nlm.nih.gov/research/cog-project/</u>
- TIGRFAM : <u>https://tigrfams.jcvi.org/cgi-bin/index.cgi</u>
- PHENO : <a href="https://phenodb.org/">https://phenodb.org/</a>

#### **Target function database**

E Select/Unselect all



16S : at least 'EC' or/and 'KO' should be chosen (EC for Metacyc pathway analysis or/and KO for KEGG pathway analysis) - others values are optionnal. ITS and 18S : 'EC' only available. (--functions)

#### NSTI cut-off

0.41

Any sequence with an NSTI above this threshold will be out. (default: 2) (--max-nsti)

#### Identity alignment cut-off

0.9

Percentage identity of the alignment between the input sequence and the PICRUSt2 reference sequence. Below this threshold, all sequences will be discarded. (default: None) (--min-blast-ident)

#### Coverage alignment cut-off



Coverage identity of the alignment between the input sequence and the PICRUSt2 reference sequence. Below this threshold, all sequences will be discarded. (default: None) (--min-blast-cov)

#### HSP method

⊘ mp	
O emp_prob	
O pic	
O scp	
O subtree_average	

Hidden-state prediction method to use: maximum parsimony (mp), empirical probabilities (emp\_prob), continuous traits prediction using subtree averaging (subtree\_average), continuous traits prediction with phylogentic independent contrast (pic), continuous traits reconstruction using squared-change parsimony (scp) (default: mp). (--hsp-method)

#### **Target function database**

Select/Unselect all



16S : at least 'EC' or/and 'KO' should be chosen (EC for Metacyc pathway analysis or/and KO for KEGG pathway analysis) - others values are optionnal. ITS and 18S : 'EC' only available. (--functions)

#### NSTI cut-off

0.41

Any sequence with an NSTI above this threshold will be out. (default: 2) (--max-nsti)

#### Identity alignment cut-off

0.9

Percentage identity of the alignment between the input sequence and the PICRUSt2 reference sequence. Below this threshold, all sequences will be discarded. (default: None) (--min-blast-ident)

#### Coverage alignment cut-off

0.9

Coverage identity of the alignment between the input sequence and the PICRUSt2 reference sequence. Below this threshold, all sequences will be discarded. (default: None) (--min-blast-cov)

#### HSP method

⊘ mp			
O emp_prob			
O pic			
O scp			
O subtree_average			

Hidden-state prediction method to use: maximum parsimony (mp), empirical probabilities (emp\_prob), continuous traits prediction using subtree averaging (subtree\_average), continuous traits prediction with phylogentic independent contrast (pic), continuous traits reconstruction using squared-change parsimony (scp) (default: mp). (--hsp-method)

#### Function table choice: Which default pre-calculated count table to use ?

- For 16S rRNA gene you can choose between: 'EC', 'KO', 'PFAM', 'COG', 'TIGRFAM', and/or 'PHENO'.
   You must select at least 'EC' or 'KO' because the information from Metacyc (EC) or KEGG (KO) are required.
- For ITS and 18S markers, 'EC' is only available.

#### **Target function database**

#### Select/Unselect all

× EC 16S : at least 'EC' or/and 'KO' should be chosen (EC for Metacyc pathway analysis or/and KO for KEGG pathway analysis) -120 NSTI = 0.41others values are optionnal. ITS and 18S : 'EC' only available. (--functions) NSTI cut-off 100 0.41 R Any sequence with an NSTI above this threshold will be out. (default: 2) (--max-nsti) 80 Identity alignment cut-off 0.9 . . 60 Percentage identity of the alignment between the input sequence and the PICRUSt2 reference sequence. Below this threshold, all sequences will be discarded. (default: None) (--min-blast-ident) 40 Coverage alignment cut-off 0.1 0.2 0.3 -0.10 0.9

Coverage identity of the alignment between the input sequence and the PICRUSt2 reference sequence. Below this threshold, all sequences will be discarded. (default: None) (--min-blast-cov)

#### HSP method

⊘ mp O emp_prob			
O pic			
O scp O subtree_average			

Hidden-state prediction method to use: maximum parsimony (mp), empirical probabilities (emp\_prob), continuous traits prediction using subtree averaging (subtree\_average), continuous traits prediction with phylogentic independent contrast (pic), continuous traits reconstruction using squared-change parsimony (scp) (default: mp). (--hsp-method)

#### **Target function database**

Select/Unselect all

<u> </u>
_

16S : at least 'EC' or/and 'KO' should be chosen (EC for Metacyc pathway analysis or/and KO for KEGG pathway analysis) - others values are optionnal. ITS and 18S : 'EC' only available. (--functions)

#### NSTI cut-off

0.41

Any sequence with an NSTI above this threshold will be out. (default: 2) (--max-nsti)

#### Identity alignment cut-off

D	.9		

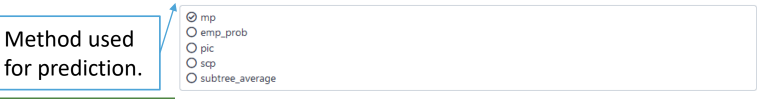
Percentage identity of the alignment between the input sequence and the PICRUSt2 reference sequence. Below this threshold, all sequences will be discarded. (default: None) (--min-blast-ident)

#### Coverage alignment cut-off

0.9

Coverage identity of the alignment between the input sequence and the PICRUSt2 reference sequence. Below this threshold, all sequences will be discarded. (default: None) (--min-blast-cov)

#### HSP method



Hidden-state prediction method to use: maximum parsimony (mp), empirical probabilities (emp\_prob), continuous traits prediction using subtree averaging (subtree\_average), continuous traits prediction with phylogentic independent contrast (pic), continuous traits reconstruction using squared-change parsimony (scp) (default: mp). (--hsp-method)

FROGSFUNC\_2\_functions: frogsfunc\_functions\_unstrat\_EC.tsv

FROGSFUNC\_2\_functions: EC\_copynumbers\_predicted.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_excluded.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_weighted\_nsti.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_marker\_norm.tsv

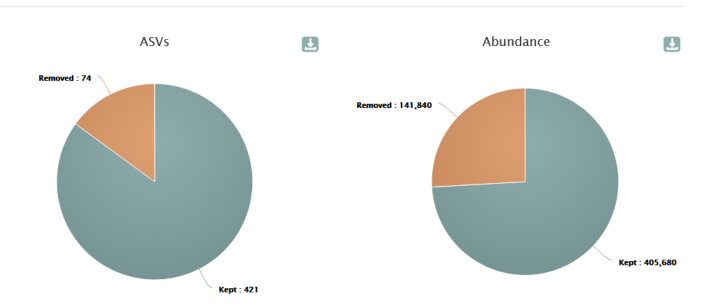
FROGSFUNC\_2\_functions: frogsfunc\_functions.fasta

FROGSFUNC\_2\_functions: frogsfunc\_functions.biom

FROGSFUNC\_2\_functions: report.html

### FROGSFUNC\_2 : report.html

ASVs are excluded if the associated NSTI is above the threshold, or if the alignment values are below the thresholds.

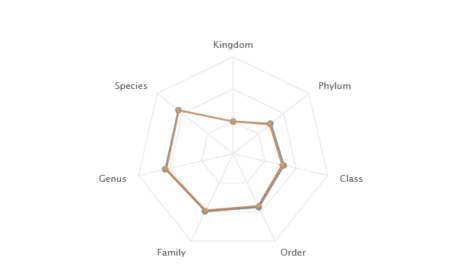


Metagenome functional profile summary

### FROGSFUNC\_2 : report.html

Number of different taxonomic ranks before (green) and after (orange) application of the filters.

Remaining diversity after filtering for functional inference



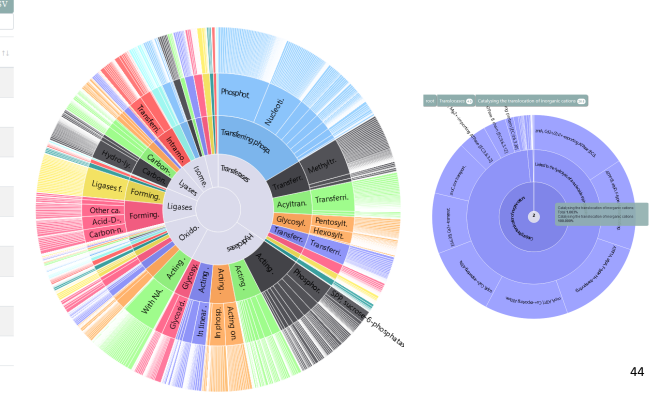
Number of different taxonomic observations per rank before and after applying NSTI or alignments thresholds

- Before
- After

### FROGSFUNC\_2 : report.html

#### Function abundances per sample 🕒 Display global distribution 📥 CSV Show 10 \$ entries Search: 1 Weighted NSTI Nb function id retrieved Samples $\checkmark$ BHT0.LOT01 0.081 2,627 $\checkmark$ BHT0.LOT03 0.075 2,701 $\checkmark$ BHT0.LOT04 0.066 2,909 $\checkmark$ BHT0.LOT05 0.081 2.836 2,815 $\checkmark$ BHT0.LOT06 0.085 BHT0.LOT07 0.082 2,787 $\checkmark$ BHT0.LOT08 0.082 2,895 $\checkmark$ 2,764 BHT0.LOT10 0.074 2,553 CDT0.LOT02 0.073 2,693 CDT0.LOT04 0.053

Gene families/function from KEGG or Metacyc databases are classified according to 4 hierarchy levels. The graph shows the proportion of each level within the selected samples.



FROGSFUNC\_2\_functions: frogsfunc\_functions\_unstrat\_EC.tsv

FROGSFUNC\_2\_functions: EC\_copynumbers\_predicted.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_excluded.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_weighted\_nsti.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_marker\_norm.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions.fasta

FROGSFUNC\_2\_functions: frogsfunc\_functions.biom

FROGSFUNC\_2\_functions: report.html

ASV abundance table without excluded ASVs (NSTI, %identity or %coverage thresholds alignment).

ASV Sequence file without excluded ASVs (NSTI, blast %identity or blast %coverage thresholds).

FROGSFUNC\_2\_functions: frogsfunc\_functions\_unstrat\_EC.tsv

FROGSFUNC\_2\_functions: EC\_copynumbers\_predicted.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_excluded.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_weighted\_nsti.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_marker\_norm.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions.fasta

FROGSFUNC\_2\_functions: frogsfunc\_functions.biom

FROGSFUNC\_2\_functions: report.html

ASV normalized abundance table

### FROGSFUNC\_2: marker\_norm.tsv

Table with normalized abundances per marker copy number from FROGSFUNC\_1 step.

Ex: cluster\_123 have two 16S copies and its abundance before this normalization in BHT0.LOT01 was 35

Cluster_124	461	0	6	6	0
Cluster_123	640	35	6	55	11
Cluster_126	511	73	7	10	33

normalized	BHT0.LOT01	BHT0.LOT03	BHT0.LOT04
Cluster_1	791.0	402.0	433.0
Cluster_10	0.0	0.0	0.0
Cluster_100	0.0	0.0	0.0
Cluster_101	0.0	0.0	0.0
Cluster_102	0.0	0.0	0.0
Cluster_103	0.0	0.0	0.0
Cluster_104	0.0	0.0	0.0
Cluster_106	0.0	0.0	0.0
Cluster_108	0.0	0.0	0.0
Cluster_109	0.0	0.0	0.0
Cluster_11	3.0	0.0	4.0
Cluster_110	0.0	0.0	0.0
Cluster_111	2.0	8.0	59.0
Cluster_112	0.0	0.0	0.0
Cluster_113	0.0	4.0	5.0
Cluster_114	0.0	0.0	0.5
Cluster_115	0.0	6.0	7.0
Cluster_117	0.0	0.0	0.0
Cluster_118	0.0	46.0	6.0
Cluster_119	0.0	0.0	0.0
Cluster_12	1333.0	188.0	127.0
Cluster_120	0.0	0.0	0.0
Cluster_121	0.0	0.0	0.0
Cluster_122	0.0	0.0	0.0
Cluster_123	17.5	3.0	27.5
Cluster_125	45.5	6.0	22.5

FROGSFUNC\_2\_functions: frogsfunc\_functions\_unstrat\_EC.tsv

FROGSFUNC\_2\_functions: EC\_copynumbers\_predicted.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_excluded.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_weighted\_nsti.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_marker\_norm.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions.fasta

FROGSFUNC\_2\_functions: frogsfunc\_functions.biom

FROGSFUNC\_2\_functions: report.html

the mean of NSTI value per sample.

sample	weighted_NSTI
BHT0.LOT01	0.0814507179687713
BHT0.LOT03	0.07523644621312382
BHT0.LOT04	0.06550232405467385
BHT0.LOT05	0.08141930786656948
BHT0.LOT06	0.08495448189855995
BHT0.LOT07	0.08161575516954905
BHT0.LOT08	0.08233567661364216

FROGSFUNC\_2\_functions: frogsfunc\_functions\_unstrat\_EC.tsv

FROGSFUNC\_2\_functions: EC\_copynumbers\_predicted.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_excluded.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_weighted\_nsti.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_marker\_norm.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions.fasta

FROGSFUNC\_2\_functions: frogsfunc\_functions.biom

FROGSFUNC\_2\_functions: report.html

### FROGSFUNC\_2: excluded.tsv

#Cluster	FROGS_taxonomy	PICRUSt2_taxonomy
Cluster_2	Bacteria; Proteobacteria; Gammaproteobacteria; Enterobac	Bacteria; Proteobacteria; Gammaproteobacter
Cluster_3	Bacteria; Firmicutes; Bacilli; Lactobacillales; Lactobacillaceae;	Bacteria; Firmicutes; Bacilli; Lactobacillales; Lacto
Cluster_7	Bacteria; Firmicutes; Bacilli; Erysipelotrichales; Erysipelotricha	Bacteria; Tenericutes; Mollicutes; Acholeplasma
Cluster_9	Bacteria; Firmicutes; Bacilli; Lactobacillales; Lactobacillaceae;	Bacteria; Firmicutes; Bacilli; Lactobacillales; Lacto
Cluster_14	Bacteria; Firmicutes; Bacilli; Lactobacillales; Lactobacillaceae;	Bacteria; Firmicutes; Bacilli; Lactobacillales; Lacto
Cluster_17	$Bacteria; Firmicutes; Bacilli; Lactobacillales; Streptococcacea {\cite{cocc}} accea $	Bacteria; Firmicutes; Bacilli; Lactobacillales; Strep

exclusion_paramater	value_parameter
min_blast_coverage	coverage = 0.819099999999999
min_blast_coverage	coverage = 0.8327
min_blast_identity,min_blast_coverage,max_nsti	identity = 0.8289,coverage = 0.711,nsti = 0.4391649999999997
min_blast_coverage	coverage = 0.8632
min_blast_coverage	coverage = 0.8327
min_blast_coverage	coverage = 0.668700000000001

Information (FROGS taxonomy, PICRUSt2 taxonomy, exclusion\_parameter, value\_parameter) on deleted ASV that are out of the cut-off values selected in this step.

FROGSFUNC\_2\_functions: frogsfunc\_functions\_unstrat\_EC.tsv

FROGSFUNC\_2\_functions: EC\_copynumbers\_predicted.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_excluded.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_weighted\_nsti.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_marker\_norm.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions.fasta

FROGSFUNC\_2\_functions: frogsfunc\_functions.biom

FROGSFUNC\_2\_functions: report.html

### FROGSFUNC\_2: EC\_copynumber\_predicted.tsv

ASV	EC:1.1.1.1	EC:1.1.1.10	EC:1.1.1	1.100	EC:1.1.1.101	EC:1.1.1.102	EC:1.1.1.103	EC:1.1.1.105
Cluster_1		2	0	2	C	0 0	0	0
Cluster_10		1	0	4	C	0 0	0	0
Cluster_100		2	0	8	C	0 0	1	0
Cluster_101		0	0	2	C	0 0	0	1
Cluster_102		0	0	3	C	0 0	0	1
Cluster_103		0	0	2	C	0 0	0	0
Cluster_104		0	0	2	C	0 0	0	0
Cluster_105		1	0	3	C	0 0	0	0
Cluster_106		1	0	4	C	0 0	0	0
Cluster_107		3	0	4	C	0 0	0	0
Cluster_108		3	0	10	C	0 0	0	0
Cluster_109		0	0	2	C	0 0	0	0
Cluster_11		5	0	5	C	0 0	1	0
Cluster_110		0	0	2	C	0 0	0	0
Cluster_111		5	0	5	C	) C	1	0
Cluster_112		2	0	1	C	0 0	0	0
Cluster_113		4	0	3	C	0 0	0	0
Cluster_114		2	0	3	C	0 0	0	0

Output table of predicted function copy numbers per ASV.

One per chosen target function database (EC, KO, PFAM, COG, TIGRFAM, PHENO).

FROGSFUNC\_2\_functions: frogsfunc\_functions\_unstrat\_EC.tsv

FROGSFUNC\_2\_functions: EC\_copynumbers\_predicted.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_excluded.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_weighted\_nsti.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions\_marker\_norm.tsv

FROGSFUNC\_2\_functions: frogsfunc\_functions.fasta

FROGSFUNC\_2\_functions: frogsfunc\_functions.biom

FROGSFUNC\_2\_functions: report.html

#### FROGSFUNC\_2: unstrat\_EC.tsv

classification	db_link	observation_name	BHT0.LOT01	BHT0.LOT03	BHT0.LOT04
Oxidoreductases;Acting on the CH-OH group of donors;With NAD+ or NADP+ as acceptor;EC:1.1.1.1	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.1	EC:1.1.1.1	13205	15364	13783
Oxidoreductases; Acting on the CH-OH group of donors; With NAD+ or NADP+ as acceptor; EC: 1.1.1.100	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.100	EC:1.1.1.100	20029	21513	23461
Oxidoreductases; Acting on the CH-OH group of donors; With NAD+ or NADP+ as acceptor; EC: 1.1.1.103	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.103	EC:1.1.1.103	177	2011	334
Oxidoreductases; Acting on the CH-OH group of donors; With NAD+ or NADP+ as acceptor; EC: 1.1.1.105	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.105	EC:1.1.1.105	0	0	0
Oxidoreductases;Acting on the CH-OH group of donors;With NAD+ or NADP+ as acceptor;EC:1.1.1.108	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.108	EC:1.1.1.108	142	194	127
Oxidoreductases; Acting on the CH-OH group of donors; With NAD+ or NADP+ as acceptor; EC: 1.1.1.11	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.11	EC:1.1.1.11	2	10	2
Oxidoreductases; Acting on the CH-OH group of donors; With NAD+ or NADP+ as acceptor; EC:1.1.1.122	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.122	EC:1.1.1.122	16	109	54
Oxidoreductases; Acting on the CH-OH group of donors; With NAD+ or NADP+ as acceptor; EC: 1.1.1.125	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.125	EC:1.1.1.125	2218	505	1656

-)

From this table of abundance it is quite possible to make statistical analyses to understand the information.

### FROGSFUNC\_3\_pathways

How it works ?

- 1. Places the ASVs into a reference phylogenetic tree and predicts of marker copy number in each ASV.
- 2. Predicts number of function copy number in each ASV and calculates functions abundances in each sample and ASV abundances according to marker copy number.
- 3. Calculates pathway abundances in each sample.

It runs only on 16S, ITS or 18S

ROGSFUNC\_1\_placeseqs\_copynumber

FROGSFUNC\_2\_functions

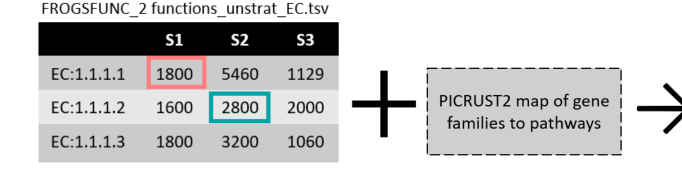
FROGSFUNC\_3\_pathways



### FROGSFUNC\_3

**FROGSFUNC\_3\_pathways** infers MetaCyc/KEGG pathway abundances based on EC/KO number abundances.

- Regroups EC/KO numbers to MetaCyc/KEGG reactions.
- Infers which MetaCyc/KEGG pathways are present based on these reactions with <u>MinPath</u>.
- Calculates and returns the abundance of pathways identified as present.



FROGSFUNC\_3 pathways\_unstrat per sample and per reference

Pathways	<b>S1</b>	S2	<b>S</b> 3
1CMET2-PWY	1289.7451	1485.2474	1233.5908
ANAEROFRUCAT-PWY	904.7455	1565.5453	1227.6231
ANAGLYCOLYSIS-PWY	1501.0804	1805.3271	1544.3206
ARG+POLYAMINE-SYN	0	49.3391	45.6559

### FROGSFUNC\_3

FROGSFUNC\_3\_pathways Calculates pathway abundances in each sample. 😭 Favorite & Versions 🗸 Options

#### **Function abundance file**

61: FROGSFUNC\_2\_functions: frogsfunc\_functions\_unstrat\_EC.tsv

TSV function abundances table from FROGSFUNC\_2\_functions tool, FROGSFUNC\_2\_functions\_unstrat\_EC.tsv for Metacyc database or FROGSFUNC\_2\_functions\_unstrat\_KO.tsv for Kegg database (unstratified table). (--input-file)

#### **Taxonomic marker**

⊘ 16S
 ○ ITS
 ○ 18S

#### Taxonomic marker of interest.

#### **Pathway reference**

O Metacyc Metacyc is the only choice for ITS and 18S

For 16S marker, choose Metacyc or KEGG in accordance with your choice in the FROGSFUNC\_1\_placeseqs\_copynumbers tool. For ITS or 18S marker, Metacyc is the only valid option.

#### Do you want to normalize the final output table ?

🕘 No

Values are divided by sum of columns, then multiplied by 10^6 (CPM values). (--normalisation)

#### **Email notification**



Send an email notification when the job completes.

### 4

normalization: values are divided by sum of columns, then multiplied by 10<sup>6</sup> (Count Per Million values).

B

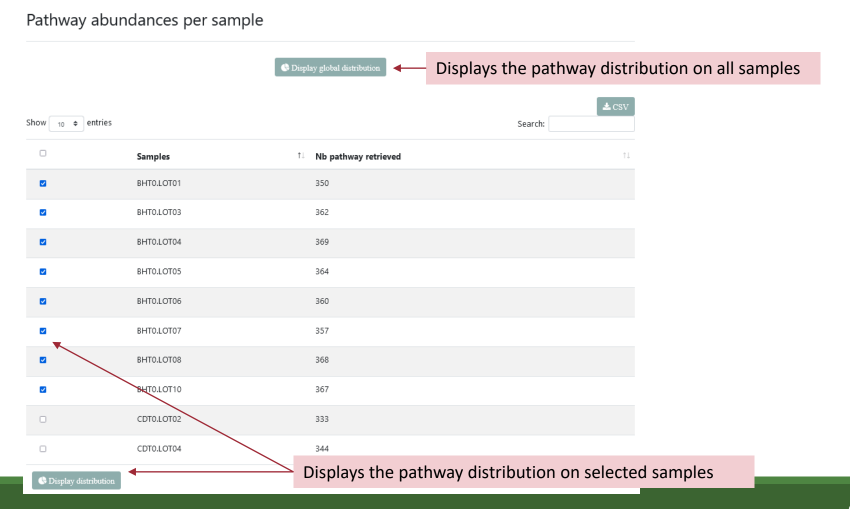
This normalization allows to compare the samples between them. But to perform more precise statistical analysis, some tools as **DESeq2 need the non-normalized abundance table** to perform the normalization by themselves.

So be careful which table to use for further analysis.

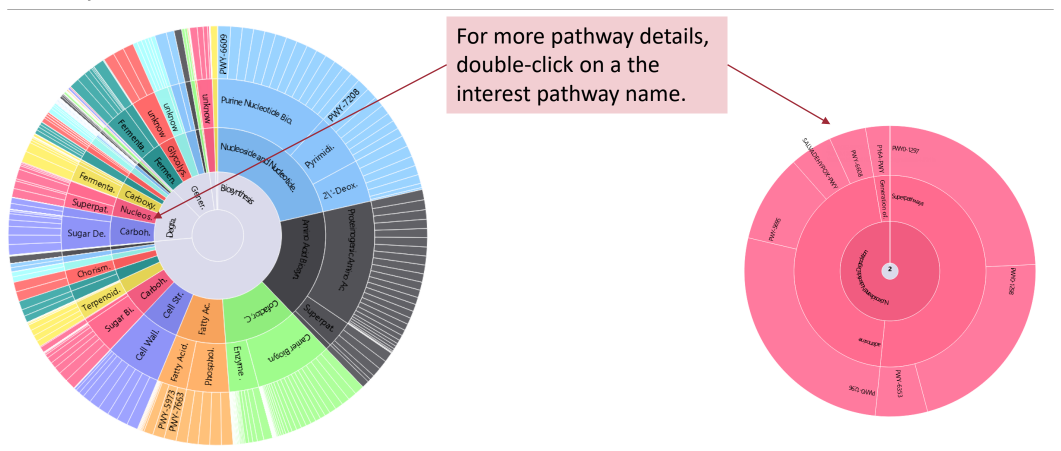
FROGSFUNC\_3\_pathways: frogsfunc\_pathways\_unstrat.tsv

FROGSFUNC\_3\_pathways: report.html

### FROGSFUNC\_3: report.html



## What is the distribution of pathway abundances in the samples ?



FROGSFUNC\_3\_pathways: frogsfunc\_pathways\_unstrat.tsv

FROGSFUNC\_3\_pathways: report.html

pathways abundance predictions of metagenome per sample.

### FROGSFUNC\_3: Pathway abundance tables



External link on the pathway

From this table of abundance it is quite possible to make statistical analyses on it to understand the information.

### **Statistics**

DIFFERENTIAL ANALYSIS

# FROGSSTAT DESeq2 preprocess

FROGSSTAT DESeq2 Preprocess import a Phyloseq object and prepare it for DESeq2 differential abundance analysis (Galaxy Version 4.1.0+galaxy1)

☆ Favorite 🚳 Versions Options

#### Type of analysis

O AS	SV .			classification		db_link	observation_name	BHT0.LOT01	BHT0.LOT03	BHT0.LOT04
Ø FU	INCTIO	NNI -		Oxidoreductases;Acting on the CH-OH group of donors;With NAD+	or NADP+ as acceptor;EC:1.1.1.1	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.1	EC:1.1.1.1	13205	15364	13783
€ FU	NCHO	л		Oxidoreductases;Acting on the CH-OH group of donors;With NAD+	or NADP+ as acceptor;EC:1.1.1.100	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.100	EC:1.1.1.100	20029	21513	23461
				Oxidoreductases;Acting on the CH-OH group of donors;With NAD+	or NADP+ as acceptor;EC:1.1.1.103	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.103	EC:1.1.1.103	177	2011	334
т			and the differential englishing ACM DECarD is may another ACM along the back	Oxidoreductases; Acting on the CH-OH group of donors; With NAD+	or NADP+ as acceptor;EC:1.1.1.105	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.105	EC:1.1.1.105	0	0	0
Type of	r data t	о репо	rm the differential analysis. ASV: DESeq2 is run on the ASV abundance table.	Oxidoreductases; Acting on the CH-OH group of donors; With NAD+	or NADP+ as acceptor;EC:1.1.1.108	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.108	EC:1.1.1.108	142	194	127
prodict	od fund	ction al	undance table from FROGSFUNC_2_function tool.	Oxidoreductases; Acting on the CH-OH group of donors; With NAD+	or NADP+ as acceptor;EC:1.1.1.11	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.11	EC:1.1.1.11	2	10	2
predicti	eu iuni	ction a	undance table non rROGSPORC_2_function tool.	Oxidoreductases;Acting on the CH-OH group of donors;With NAD+	or NADP+ as acceptor;EC:1.1.1.122	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.122	EC:1.1.1.122	16	109	54
				Oxidoreductases;Acting on the CH-OH group of donors;With NAD+	or NADP+ as acceptor;EC:1.1.1.125	https://www.genome.jp/dbget-bin/www_bget?EC:1.1.1.125	EC:1.1.1.125	2218	505	1656
Funct	tion ak	oundar	ces file							
C	C		61: FROGSFUNC_2_functions: frogsfunc_functions_unstrat_EC.tsv	-						

U U

61: FROGSFUNC\_2\_functions: frogsfunc\_functions\_unstrat\_EC.tsv

Input file of predicted function abundances (frogsfunc\_functions\_unstrat.tsv from FROGSFUNC\_2\_function tool).

#### Metadata associated to samples (format: TSV)

2: metadata_chaillou.tsv		•		
The file must contain the metadata that characterise each sample. (samplefile)		EnvType	Description	FoodType
Even and the second at the sec	BHT0.LOT01	BoeufHache	LOT1	Meat
Experimental variable	BHT0.LOT03	BoeufHache	LOT3	Meat
EnvType	BHT0.LOT04	BoeufHache	LOT4	Meat

The factor that could have an effect on ASV/FUNCTION abundances. Ex: Treatment, etc.

#### Do you want to correct a confounding factor?

False -If yes, specify the counfouding factor **Email notification** No Send an email notification when the job completes.

#### Outputs

FROGSSTAT DESeq2 Preprocess: function\_data.Rdata

FROGSSTAT DESeq2 Preprocess: function\_dds.Rdata

# FROGSSTAT DESeq2 visualisation

FROGSSTAT DESeq2 Visualisation to extract and visualise differentially abundant ASVs or functions	☆ Favorite	& Vers
(Galaxy Version 4.1.0+ galaxy1)	A ravonic	the read

Type of analysis



Type of data to perform the differential analysis. ASV: DESeq2 is run on the ASV abundance table. FUNCTION: DESeq2 is run on predicted function abundance table from FROGSFUNC\_2\_function tool.

Data object (format: data.RData)

For ASV: asv\_data.Rdata from FROGSSTAT\_Phyloseq\_Import\_Data tool - For FUNCTION: function\_data.Rdata from FROGSSTAT\_DESeq2\_Preprocess tool. (--abundanceData)

DESeq2 object (format: dds.RData)

C 73: FROGSSTAT DESeq2 Preprocess: function_dds.Rdata	dds.Rdata	•	B	
---	-----------	---	---	--

This is the result of FROGSSTAT\_DESeq2\_Preprocess tool asv\_dds.Rdata or function\_dds.Rdata (--dds)

#### **Experimental variable**

EnvType		
The factor that could have a	n effect on ASV/FUNCTION abundances. Ex : Treatment (var)	-

		EnvType	Description	FoodType
The experimental variable is it quantitative or qualitative?	BHT0.LOT01	BoeufHache	LOT1	Meat
Qualitative	BHT0.LOT03	BoeufHache	LOT3	Meat
If qualitative, choose 2 conditions to compare	BHT0.LOT04	BoeufHache	LOT4	Meat
Condition 1 considered as reference	VHT0.LOT07	VeauHache	LOT7	Meat
BoeufHache	VHT0.LOT08	VeauHache	LOT8	Meat
One condition of the experimental variable (e.g. with) (mod2)	VHT0.LOT10	VeauHache	LOT10	Meat
Condition 2 to be compared to the reference	SFT0.LOT01	SaumonFume	LOT1	Seafood
VeauHache	SFT0.LOT02	SaumonFume	LOT2	Seafood
	SFT0.LOT03	SaumonFume	LOT3	Seafood
A weather and data we falls and a local a local a local the south (south falls)				

Another condition of the experimental variable (e.g. without) (--mod1)

Adjusted p-value threshold

0.05

#### For the moment, we keep default p-value

Threshold used for statistical significance of the differentially abundant ASV/FUNCTION analysis (--padj)

FROGSSTAT DESeq2 Visualisation: ipath\_under.tsv

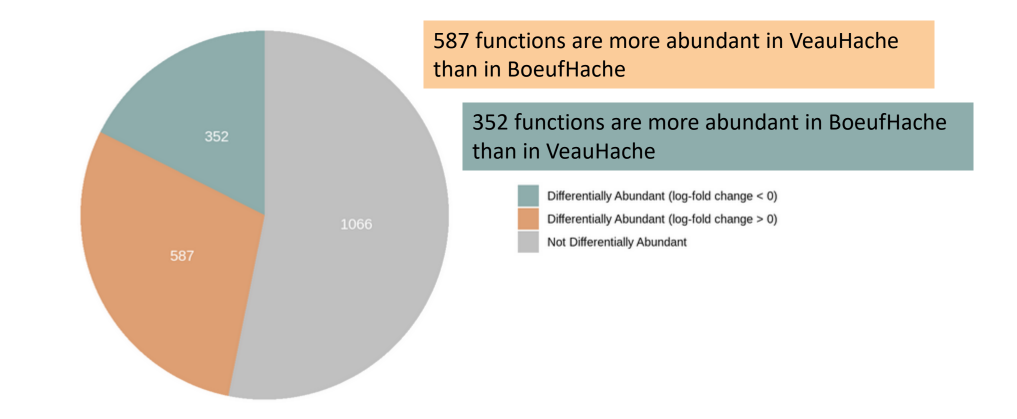
FROGSSTAT DESeq2 Visualisation: ipath\_over.tsv

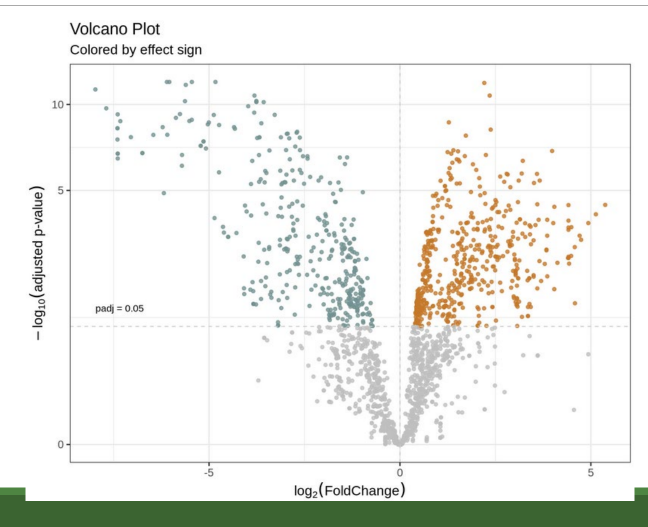
FROGSSTAT DESeq2 Visualisation: report.nb.html

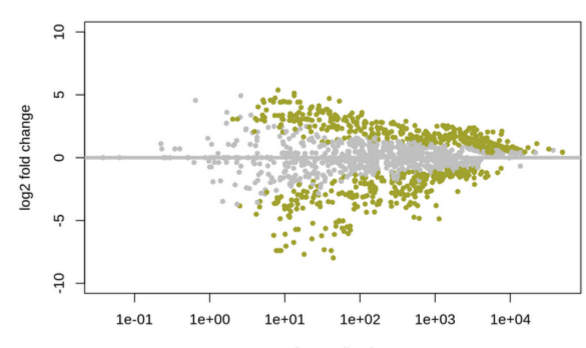
	ID 🔶		baseMean 🔺	log2FoldChange	Ifc SE 🔶	stat 🛊	pvalue 🕴	padj 🖨	Level_4	
	AI		All	All			A	,	All	
413	EC:3.4.24.30		1.96849	4.53492	1.30759	3.46814	0.000524070	0.00249661	Hydrolases	
691	EC:6.3.2.43		2.57016	-6.11515	2.29021	-2.67013	0.00758220	0.0215825	Ligases	

You chose to compare VeauHache to the reference modality BoeufHache. This implies that a positive log2FoldChange means more abundant in VeauHache than in BoeufHache.

Pie chart to view ASVs or FUNCTIONs number of Differential Abundance test

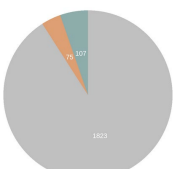


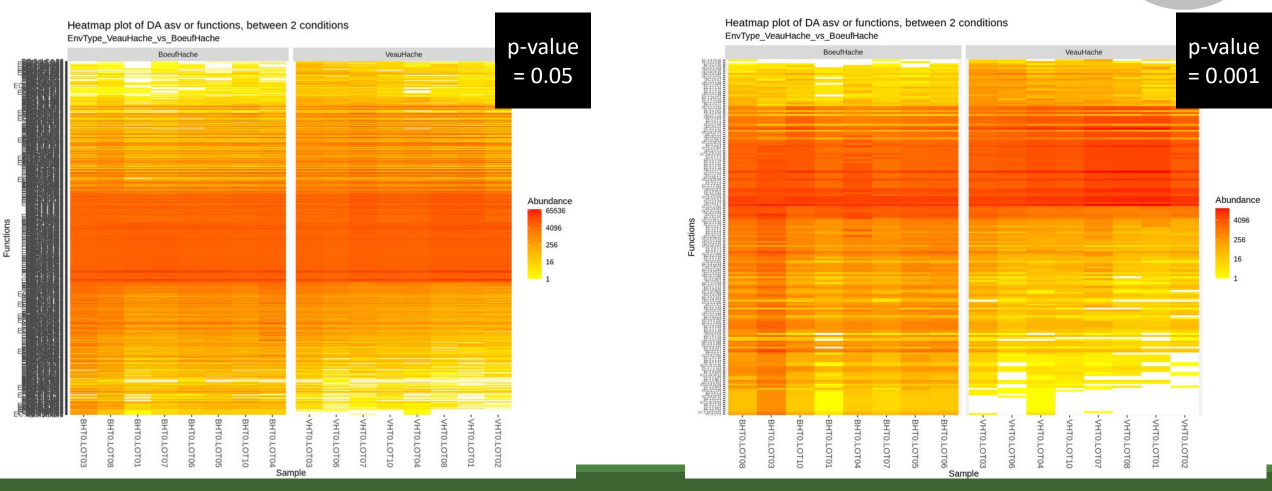




Post Normalisation DESeq2: MA plot of log2FoldChange

mean of normalized counts



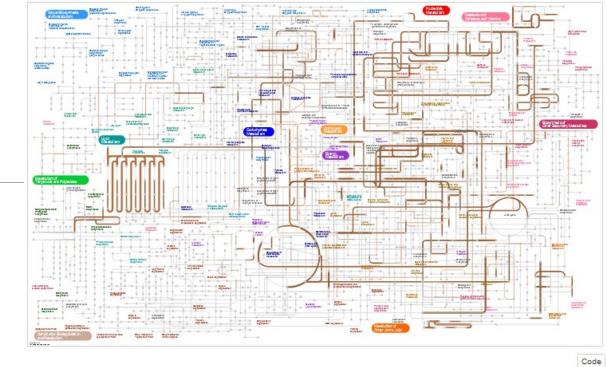


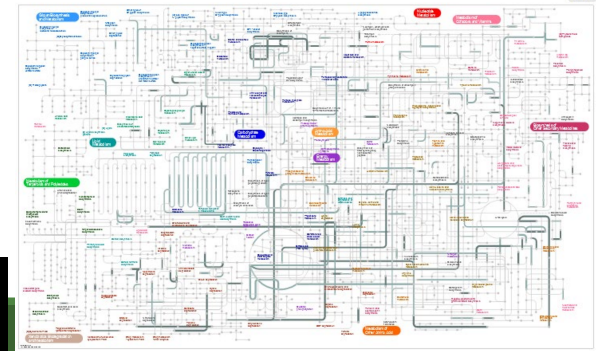
You chose to compare VeauHache to the reference modality BoeufHache. This implies that the overabundants pathways (first image) and underabundant pathways (second image) are involved in VeauHache condition.

The darker the path, the higher the log2 fold change (<1, <2, or >2).

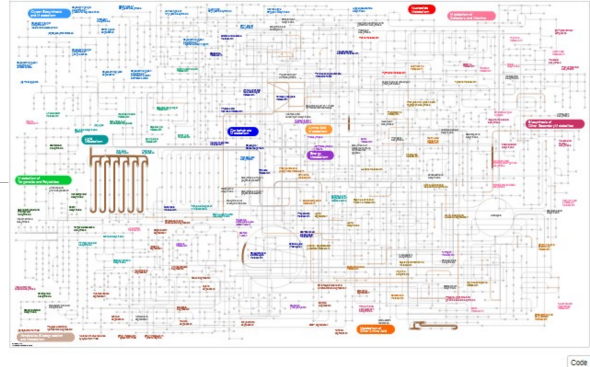
p-value

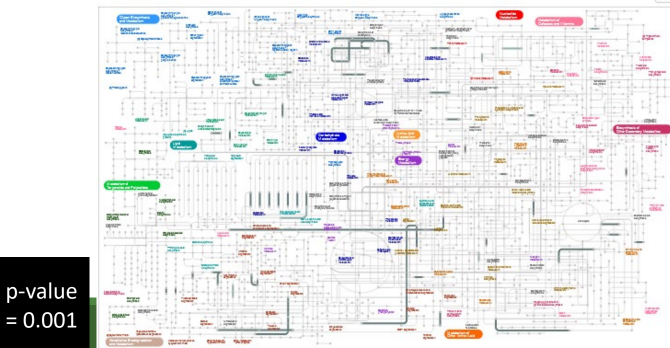
= 0.05





= 0.001





FROGSSTAT DESeq2 Visualisation: ipath\_under.tsv

FROGSSTAT DESeq2 Visualisation: ipath\_over.tsv

FROGSSTAT DESeq2 Visualisation: report.nb.html

To visualise and explore metabolic pathways with <u>IPATH3 website</u>

### Go to IPATH3

ID Color Width	
EC2.6.1.1 #637978	W12
EC2.7.8.8 #637978	W12
EC2.2.1.1 #8EADAC	W12
EC4.1.1.65 #637978	W12
EC4.1.99.3 #8EADAC	W12
EC2.2.1.2 #8EADAC	W12
EC4.2.1.118 #637978	W12
EC2.2.1.7 #8EADAC	W12
EC4.99.1.1 #8EADAC	W12
EC2.5.1.17 #8EADAC	W12

#### ID Color Width

504 3 00 4 #D500 45 1440

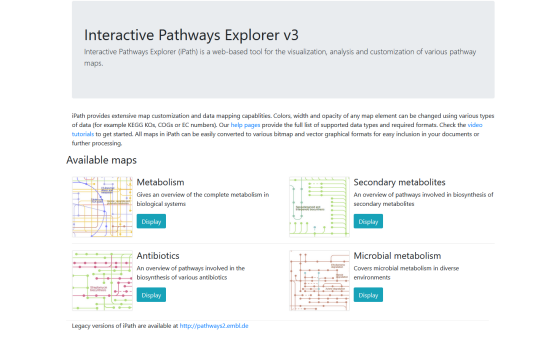
EC1.3.99.4 #DE894E W12
EC2.7.1.175 #DE894E W12
EC3.4.25.1 #DE894E W12
EC1.3.99.5 #DE894E W12
EC3.3.2.8 #DE894E W12
EC2.7.7.53 #DE894E W12
EC2.5.1.68 #DE894E W12
EC1.14.14.12 #DE894E W12
EC2.1.1.219 #DE894E W12
EC2.1.1.220 #DE894E W12

ipath\_over.tsv

#### ipath\_under.tsv

#### https://pathways.embl.de/

PATH 3 Pathway maps Tools Share Help \*



Create a free account and paste "over" or "under" data

