

### FROGSFUNC: Smart integration of PICRUSt2 software into FROGS pipeline

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## **FROGSFUNC:**

# Smart integration of PICRUSt2 software into FROGS pipeline

Vincent DARBOT<sup>1</sup>, Moussa SAMB<sup>1</sup>, Maria BERNARD<sup>2</sup>, Olivier RUÉ<sup>3</sup> and Géraldine PASCAL<sup>1</sup>

<sup>1</sup> GenPhySE, Université de Toulouse, INRAE, ENVT, F-31326, Castanet Tolosan, France. <sup>2</sup> Univ. Paris-Saclay, INRAE, AgroParisTech, GABI, SIGENAE, F-78352, Jouy-en-Josas, France.<sup>3</sup> Univ. Paris-Saclay, INRAE, BioinfOmics, MIGALE bioinformatics facility, Jouy-Corresponding Author: vincent.darbot@inrae.fr en-Josas, France

















## Background



Metabarcoding is the large-scale taxonomic identification of complex environmental samples via analysis of DNA reads of one marker gene (16S, ITS, 18S, COI, etc.).

The aim of metabarcoding analysis is to provide a table of abundance of OTUs/ASVs, as close as possible to the species, per sample as well as a descriptive statistical analysis of the composition of the targeted a known sequence having that function. microbial population of the samples.

What metabolic functions are presents in a microbial community ?

Functional inference: we assume that an organism has a metabolic function by the presence in the organism's genome of



The various tools developed within FROGS <sup>[1,2]</sup> offers such results. They allow users to process their data in command lines or *via* in a user-friendly Galaxy<sup>[3]</sup> interface and to obtain different graphical and descriptive outputs.

Unlike metagenomics<sup>[4]</sup>, metabarcoding does not provide these functional profiles of a microbial population, by being restricted to one marker gene.

PICRUSt2<sup>[5]</sup> bypass this restriction and obtain a prediction of the functional potential of a sample, at low cost.

Firstly, PICRUSt2 placed the marker gene (16S, ITS or 18S) sequences of interest into a its reference tree, that is used as the basis of functional predictions. After, it predicts number of marker and function copy number in each OTU. Then, for each sample, it calculates functions abundances and finally, pathway abundances are inferred, based on functional profile.



## **Functional profiling within FROGS**

fasta sequences sample1 sample2

Workflow

**Ability to use Galaxy interfaces** 





Milatonia - obour	
•	FROGSFUNC_step3_functions Calculates functions abundances in each sample. (Galaxy Version 40.0+galaxy1)
blom). (~input-blom)	
• 8	Biom Tile
ree.nwk(. (-tree)	D D 70: FROGSFUNC_step1_placeseqs.frogsfunc_placeseqs.biom
	The abundance file i.e. FROGSFUNC_step1_placeseqs tool output file (frogsfunc_placeseqs.biom). (input-biom) Function file
	Image: Comparison of the second se
	Copy number table of functions present in the predicted genome for each OTU i.e. FROGSFUNC_step2_copynumbers tool output file (frogsfunc_copynumbers_predicted_functions.tsv). (function)
pathway analysis) - others values are optionnal. ITS and 185.° EC only available.	Marker file       Image: Discrete black     T3: FROGSFUNC_step2_copynumbers: frogsfunc_copynumbers_marker.tsv
	Table of predicted marker copy number i.e. FROGSFUNC_step2_copynumbers output (frogsfunc_copynumbers_marker.tsv). (marker)
), continuous traits prediction using subtree averaging (subtree, average), continuous traits prediction with phylogentic independent contrast (pic), continuous traits reconstruction using	0.36
	Any sequence with an NSTI above this threshold will be out. (default: 2) (max-nsti)
	Email notification
	No No
	Send an email notification when the job completes.
	✓ Execute

### Various tables and graphical outputs are displayed to make the experience intuitive



FROGS highlights important information that are very useful for data interpretation. Some "hidden" PICRUSt2 outputs are exploited: incongruence reporting between taxonomic affiliations, Nearest Sequenced Taxon Index (NSTI) threshold confidence indicator, decision support graphic to help choosing the NSTI threshold





RESALAB





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