



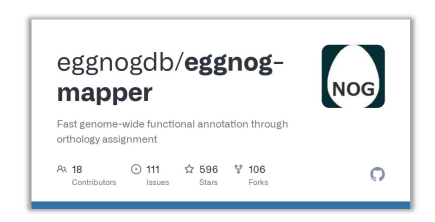
## **KEGGaNOG: A Streamlined Solution** for KEGG Pathway Annotation



Ilia Popov<sup>1</sup>, Igor Popov<sup>1</sup>, Alexey Ermakov<sup>1</sup>

<sup>1</sup> «Bioengineering and Veterinary Medicine» Faculty, Don State Technical University, Rostov-on-Don, Russia

## Introduction & Purposes of use



eggNOG-mapper is a tool for fast functional annotation of novel sequences

- It uses **precomputed orthologous groups** and phylogenies from the eggNOG database
- Its annotations are well detailed
- However, its **hard to interpret** them
- For instance, it is **hard to understand** which KEGG Orthology (KO) terms are coded in 'KEGG\_ko' column of eggNOG-mapper annotations!

### ??? KEGG\_ko K00002 K13577 K01078 K22390 K19356 K22032

K08139 K08141 K20523 K08504

K01876 K22503

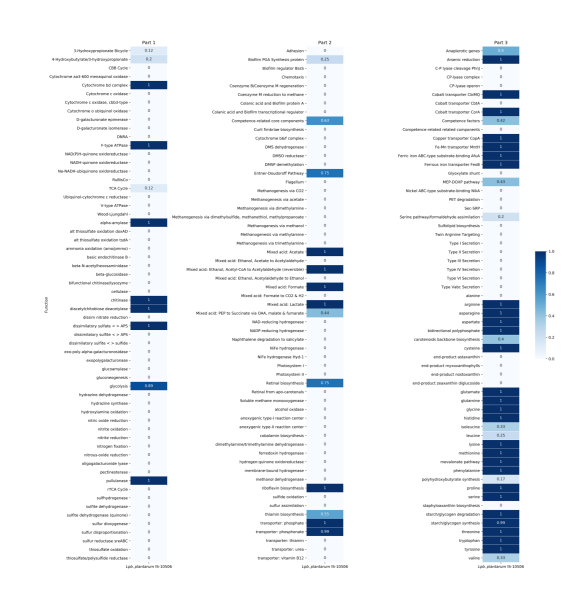
### **KEGGaNOG** solves this problem:

- It takes eggNOG-mapper annotations
- Extracts 'KEGG\_ko' column
- Decodes KO terms with KEGG-Decoder and creates .tsv file with pathways completeness
- Provides several visualization options

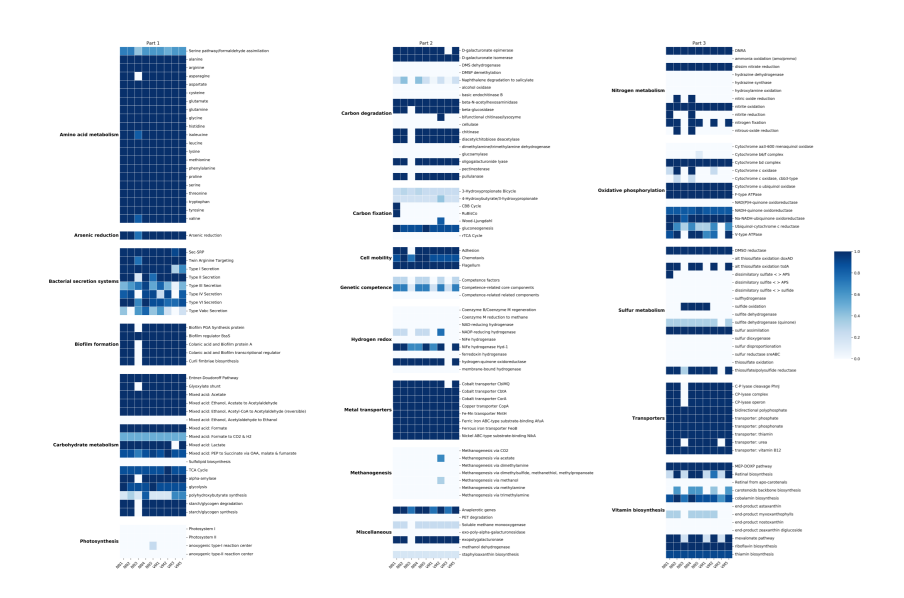
#### **KEGGaNOG** is perfect for

- Annotating bacterial metabolic profiles (e.g. probiotics)
- Annotating metagenomes metabolic profiles

## Sample outputs



Complete metabolic profile of probiotic Lpb. plantarum IS-12506. KEGGaNOG's default visualization with single sample layout. Heatmap depicts pathways completeness (0 to 1).



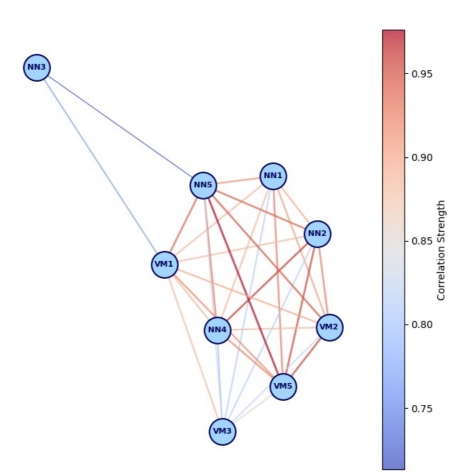
Complete metabolic profiles of 10 metagenome samples. KEGGaNOG's default visualization with multi sample layout.

# Acknowledgements

The work was carried out under the Russian Science Foundation grant № 23-14-00316

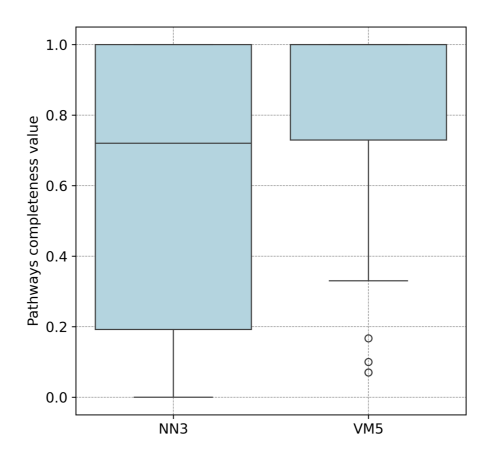
## **User APIs**

User is also provided with several other options of visualization



### **Option 1: Correlation network**

**KEGGaNOG** builds correlation matrix and plots it as the network. User is provided with options to adjust the plot (cmap, minimal correlation to plot etc.) This function allows to compare samples between each other and see the pattern – which sample differs from others



### **Option 2: Boxplots**

KEGGaNOG uses the pathways completeness data and compares the completeness tendency among several samples This function allows to compare samples in terms of which sample has core complex metabolic profile and which - not.

Several other APIs are under development at the moment:

- Spider-plot API to compare the completeness of one specific pathway between samples
- 2. PCoA API to visualize metabolic profile similarity and divergence, highlighting clustering patterns and functional shifts

Maintenance updates are also under active development

## Instructions



GitHub and README



Python Package Index