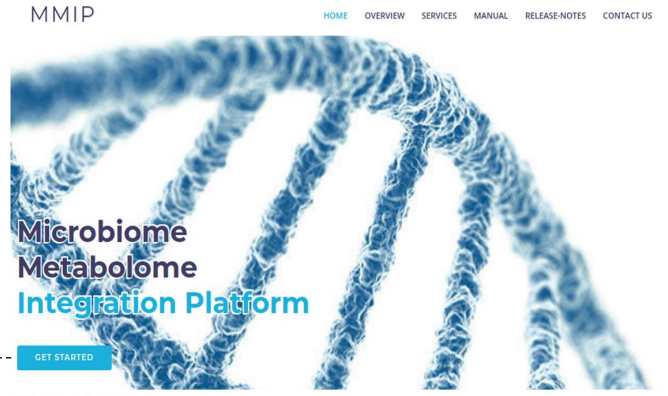
#### **WORKING MANUAL OF MMIP**

#### **Notes:**

- 1. # Red arrow and number box indicates additional functionality.
- **2.** -----▶ Black dotted arrow indicates feature information.
- **3.** All the comparison and the analyses are between two groups (not more).
- **4.** By the term "significant", statistical significance is referred to.
- **5.** All the analysis diagrams are made using Plotly and will feature the following tools (as shown below) on the top right corner of each diagram, allowing the user to access features like zoom, edit, download and other interesting tools.





# Overview

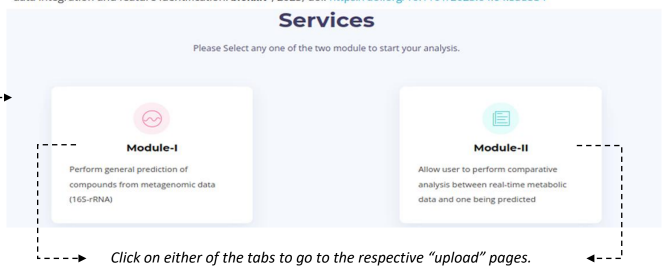
Click to use the respective services

Welcome to MMIP Server.

This server is designed for prediction of the metabolites that could be produced by the microbial community during different health condition by using their metagenomics data (16S-rRNA). Furthermore, it also helps you to compare your real-time metabolomic (untargeted) data with predicted metabolite and helps to see what could be the most probable source of the metabolites.

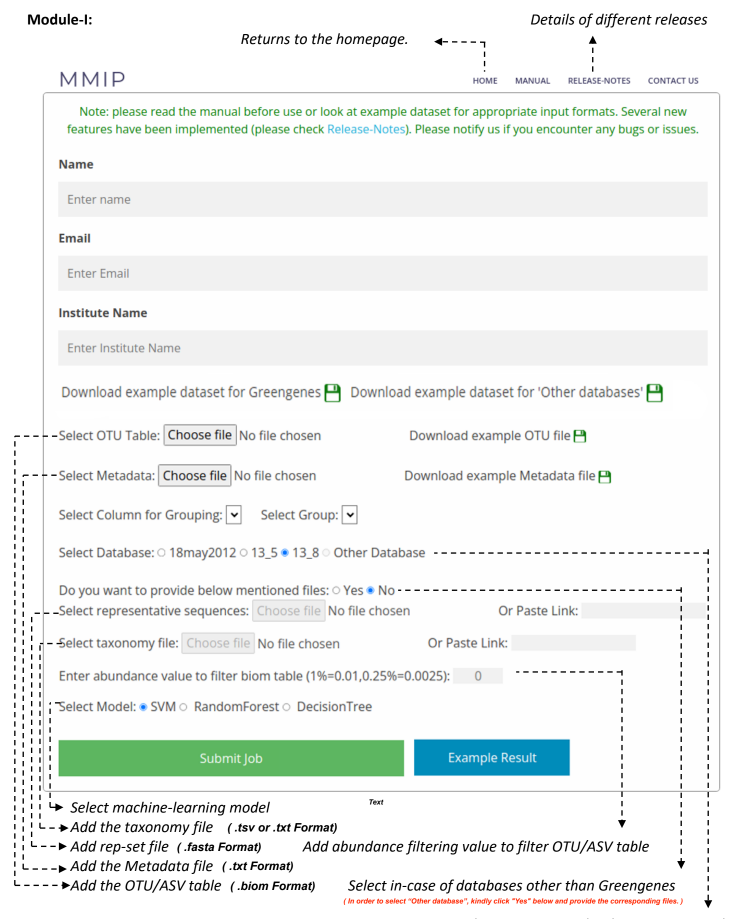
- Helps in predicting metabolites from the metagenomic data.
- ⊗ Helps in finding the most probable source of the metabolite (microbe responsible for the production).
- Predict important feature at OTU, Compound, Enzyme level by machine learning approaches.
- Helps in interlinking important feature.

**You can access preprint here:** Anupam Gautam, Debaleena Bhowmik, Sayantani Basu, Abhishake Lahiri, Wenhuan Zeng, Sandip Paul. Microbiome Metabolome Integration Platform (MMIP): a web-based platform for microbiome and metabolome data integration and feature identification. *bioRxiv*, 2023; doi: https://doi.org/10.1101/2023.04.04.535534



(Fig. 1)

## **UPLOAD PAGES**



Select Greengenes database version used

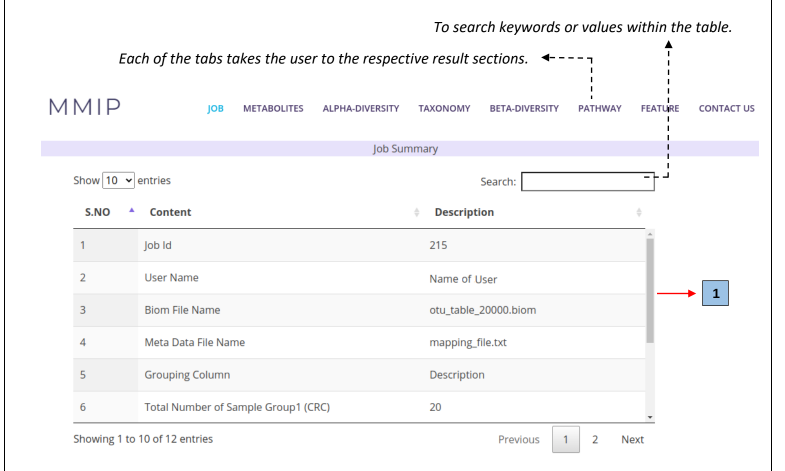
# Module-II:

MMIP	HOME	MANUAL	RELEASE-NOTES	CONTACT US				
Note: please read the manual before use or look at example dataset features have been implemented (please check Release-Notes). Please								
Name								
Enter name								
Email								
Enter Email								
Institute Name								
Enter Institute Name								
Download example dataset for Greengenes 💾 Download exam	nple data	set for 'C	ther databas	ses' 💾				
Select OTU Table: Choose file No file chosen Download	example	OTU file [	3					
Select Metadata: Choose file No file chosen Download	example N	Metadata	file 💾					
Select Column for Grouping: Select Group:								
Select Metabolomics Feature table (optional): Choose file No file chose	en	Lo	ook at the exa	mple file 💾				
Select Database: ○ 18may2012 ○ 13_5 ● 13_8 ○ Other Database								
Do you want to provide below mentioned files: O Yes  No Select representative sequences: Choose file No file chosen	Or P	aste Link:						
Select taxonomy file: Choose file No file chosen Or Past	e Link:							
Enter abundance value to filter biom table (1%=0.01,0.25%=0.0025):	0							
Select Model: ● SVM ○ RandomForest ○ DecisionTree								
Submit Job Exa	mple Res	ult						

└ - - ► Add metabolite table from metabolomic processing (.txt Format)

#### **RESULT PAGES**

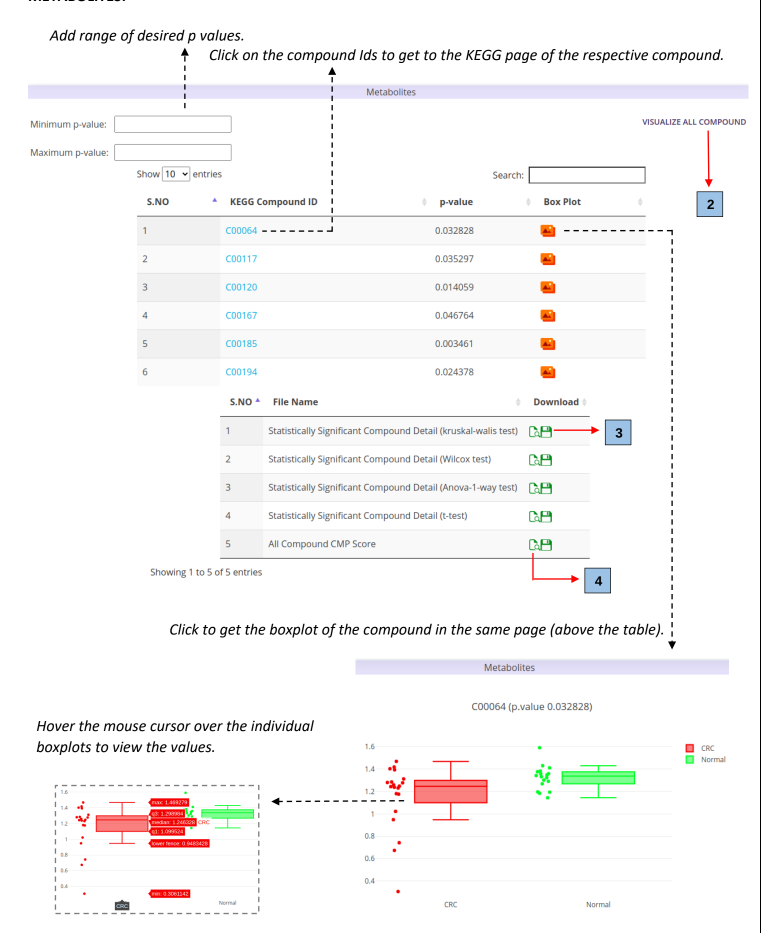
#### **USER AND JOB RELATED INFORMATION:**



(Fig. 4)

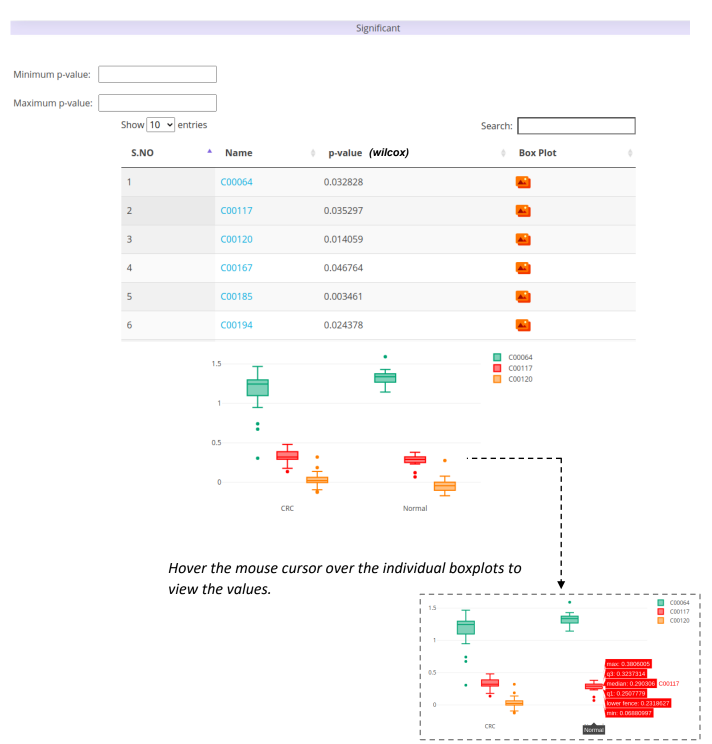
Scroll the table for additional information including number of samples in group 2, Institute name, email id, start date, time and status of the job.

#### **METABOLITES:**



(Fig. 5)

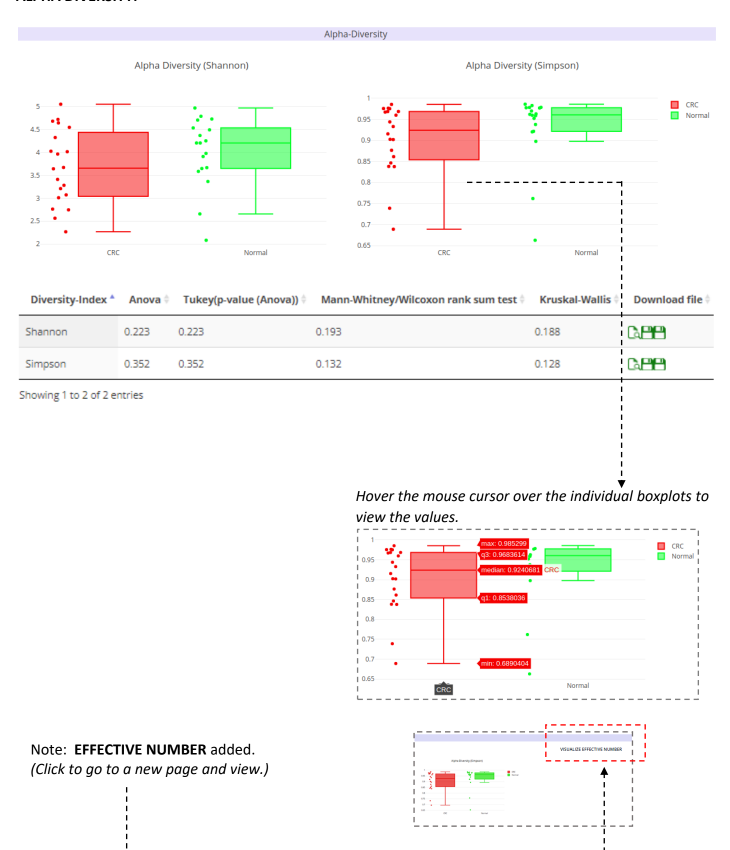
Click the tab to view all the compounds and more. The following page will open with the significant compounds. The boxplot shows the median values and differences between the two groups.



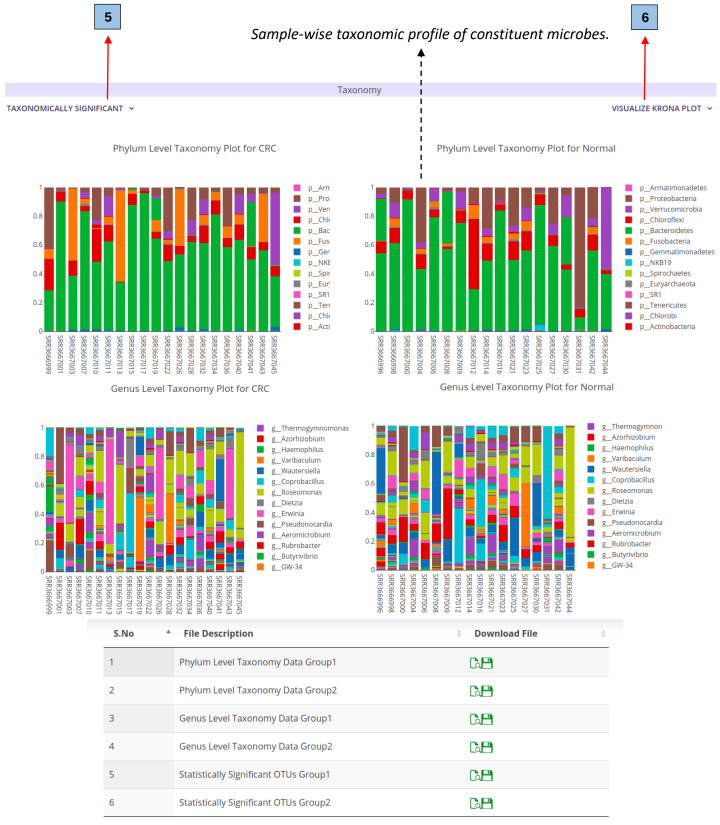
(Fig. 6)

- **3** Click to download the data table (tsv format).
- 4 Click to open the data table in a new webpage (tsv format).

## **ALPHA DIVERSITY:**

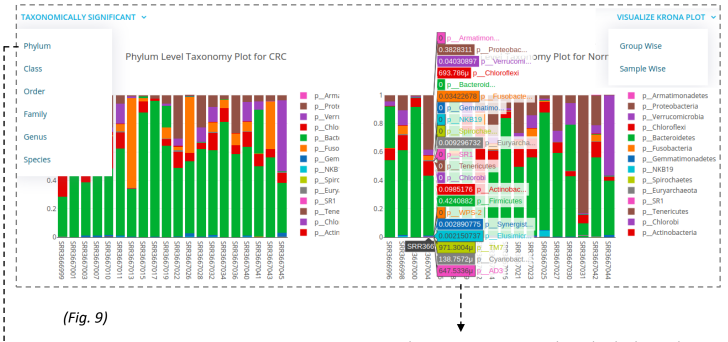


#### **TAXONOMIC DIVERSITY:**



Showing 1 to 6 of 6 entries

Click on this tab to find a drop-down list (shown below), from which user can select the desired taxonomic level to check significant ones and is visualized in a new webpage, also depicted below.

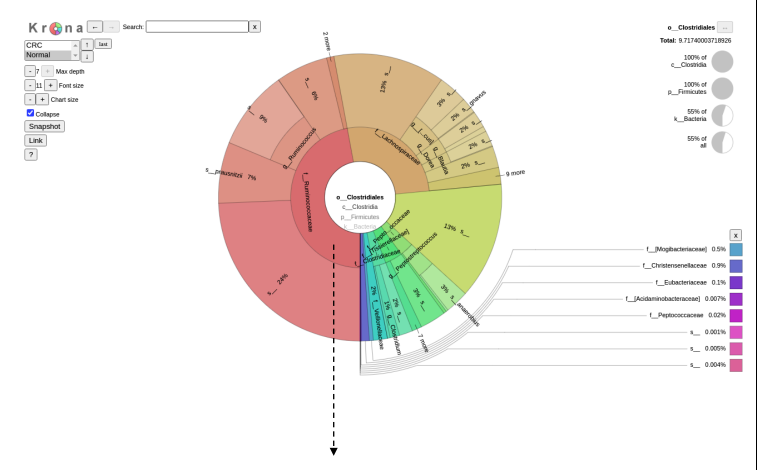


Click on one of the taxonomic level to go to the following page showing the taxa that are significant. Hover the mouse cursor over the individual samples to view the different taxa and their relative abundance.

			Significant phylum			
_						
Minimum p-value:						
Maximum p-value:						
	Show 10 ✔ ent	tries	Search:			
	S.NO	^ Name	p-value (wilcox)	Box Plot		
	1	p_Fusobacteria	0.033734	<u></u>		
	Showing 1 to 1 of 1 entries			Previous 1 Next		
		0.6		p_Fusobacteria		
		0.5				
		0.4				
		0.3				
		0.2				
		0.1				
		0 CRC	Normal			
		CKC	NOTITIAL			

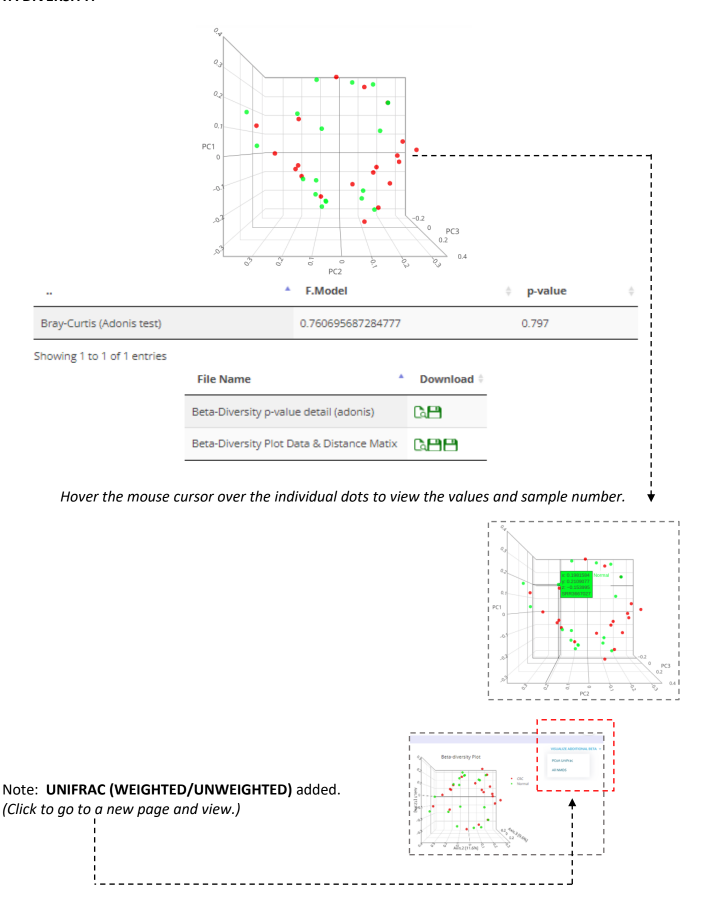
(Fig. 10)

Click on this tab to open the Krona plots representing the taxonomic abundance profile, both groupwise and sample-wise (as shown in fig. 9). This is an interactive plot, and is another way of depicting the taxonomic profiles.

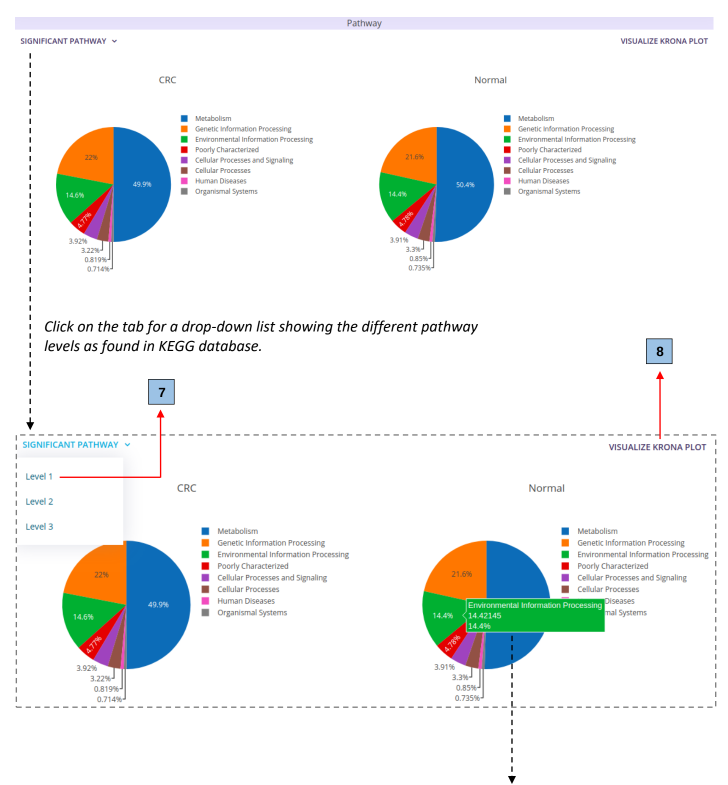


Click on the different segments to see its constituent taxa and at different levels. The ring can be collapsed back into its higher levels of taxa.

## **BETA DIVERSITY:**

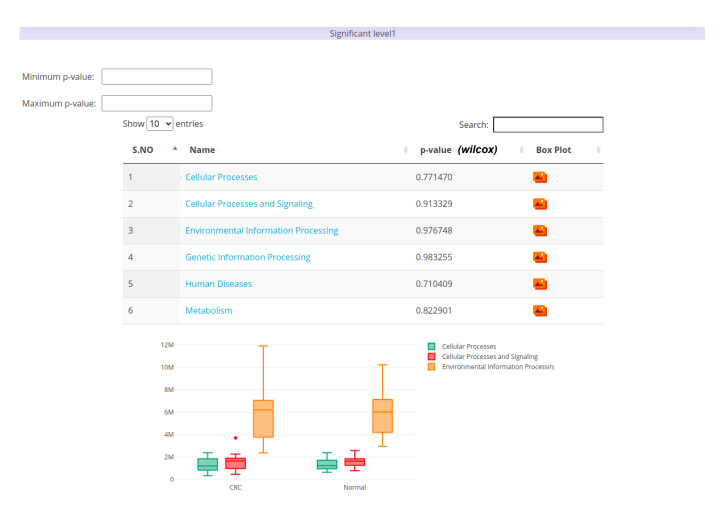


#### **PATHWAY ANALYSIS:**



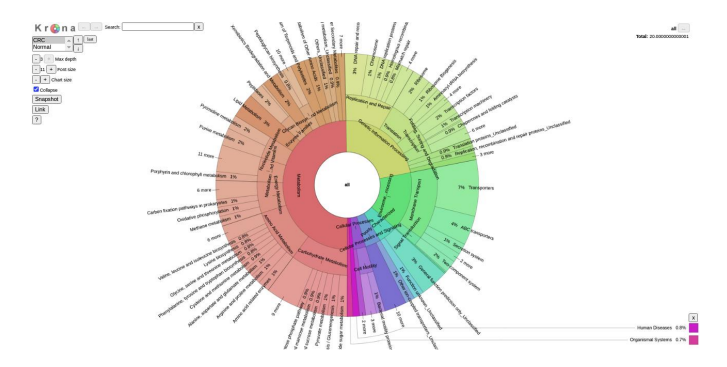
Hover the mouse cursor over the different section to view the pathway name and the percentage of its contribution to overall metabolism in that group.

7 Click on one of the levels to check the significant pathways in that level and it opens in a new page.



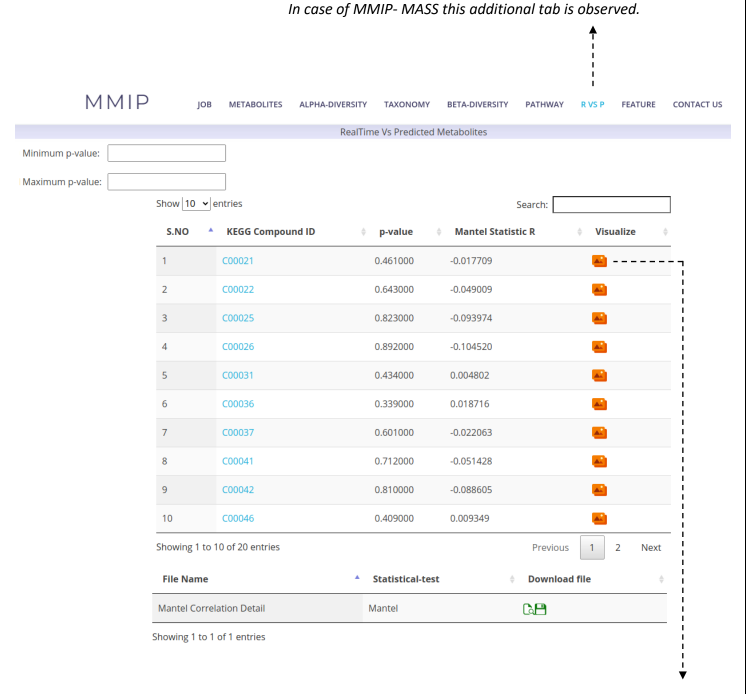
(Fig. 14)

8 Click on this tab to visualize the Krona plot for metabolic pathway abundances between two groups.



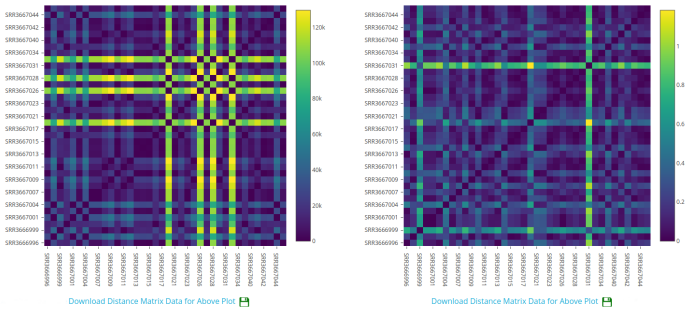
(Fig. 15)

## **REAL TIME V/S PREDICTED METABOLITES:**

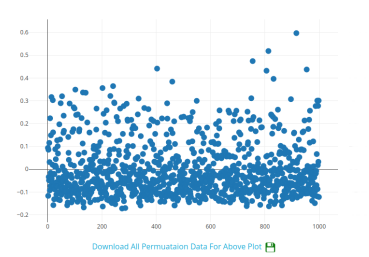


Click to open the correlation results of this metabolite in a new page (as shown in fig 17).

Distance Matrix For Predicted Time Metabolite (C00021 Method=Euclidean)



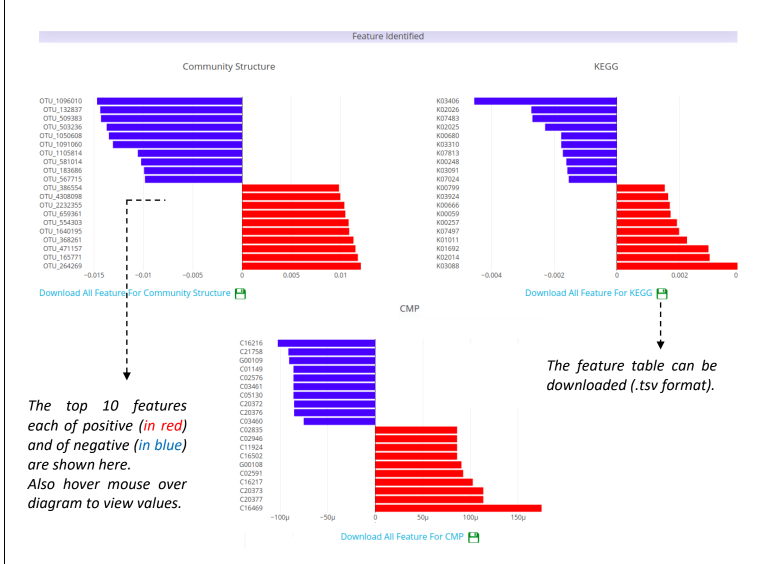
Mantel Test (Pvalue= 0.461, R-Statistics=-0.0177093134075238)

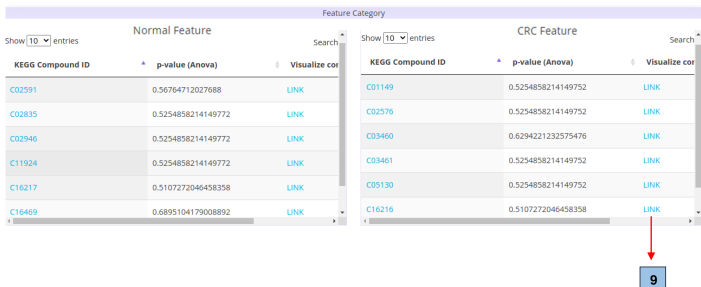


## **FEATURE PREDICTION:**



ROC curve and evaluation metrics of the selected machine learning model shown.





(Fig. 19)

9 Click on the link to visualize the following page which shows the predicted microbial source of metabolites (predicted as feature by our algorithm), in a tree as well as tabular form. [all p values provided in adjacent brackets]

