

Taxonomic and Bioactivity Characterizations of Mameliella alba Strain LZ-28 Isolated from Highly Toxic Marine Dinoflagellate *Alexandrium catenella* LZT09

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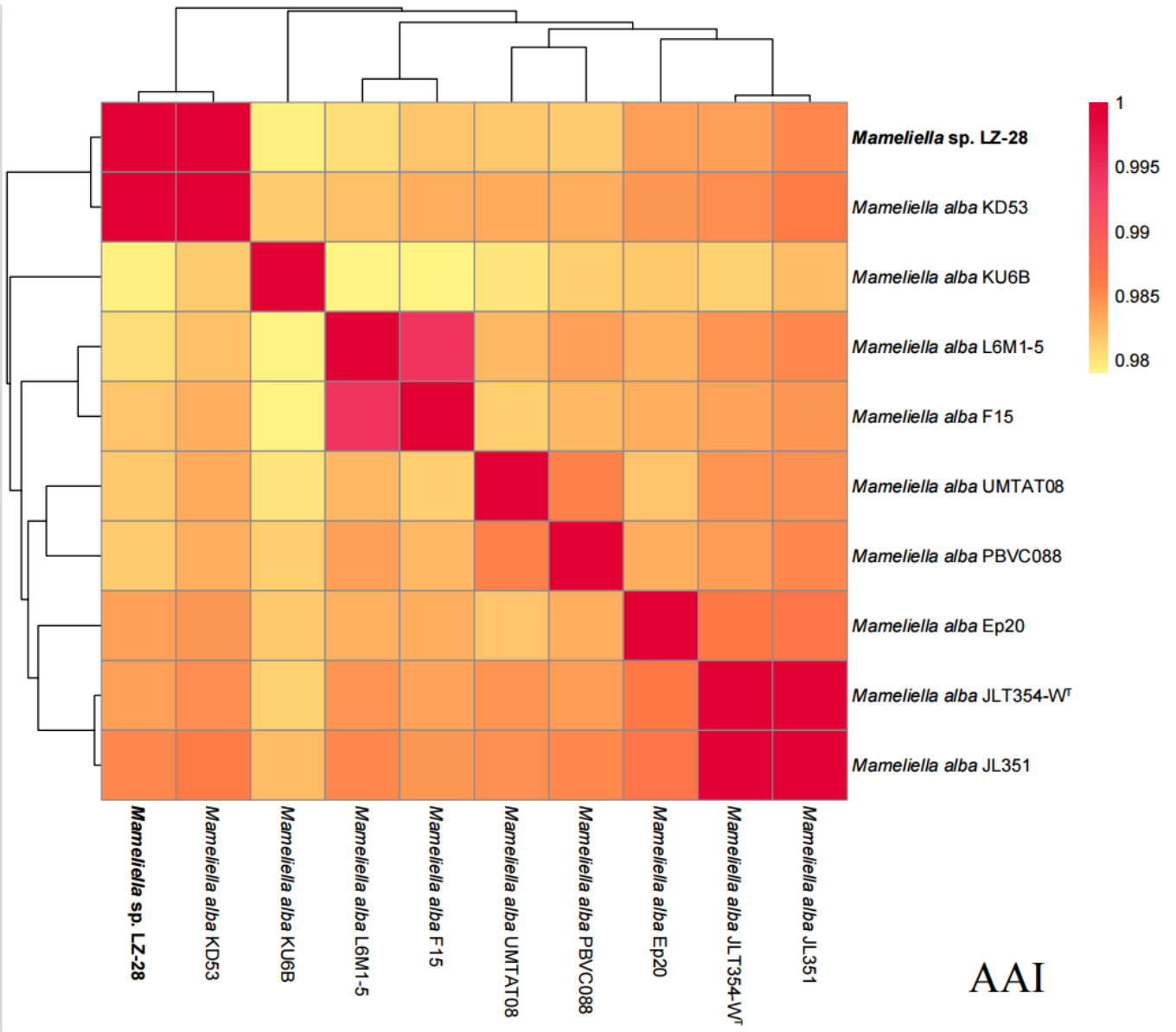
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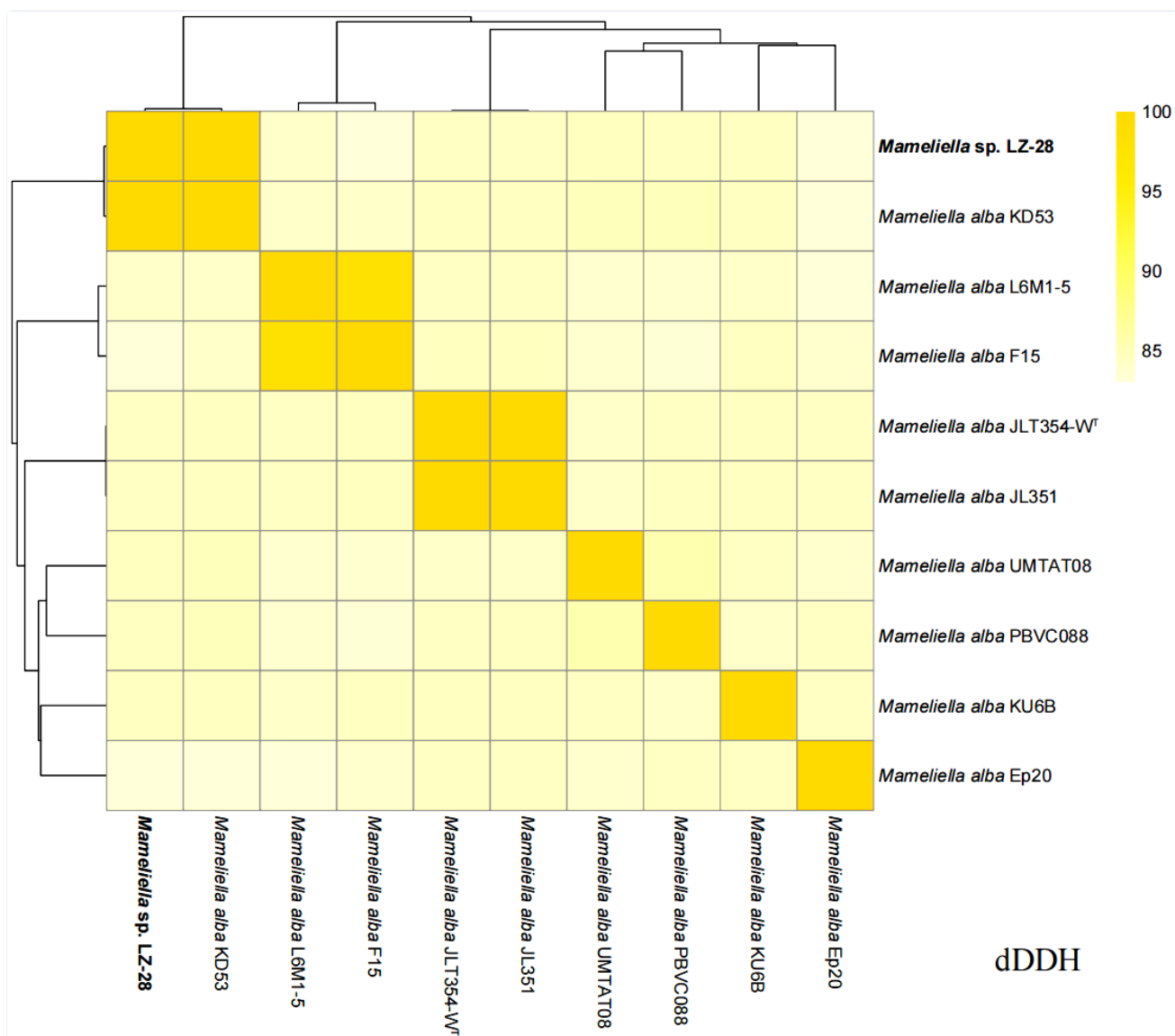
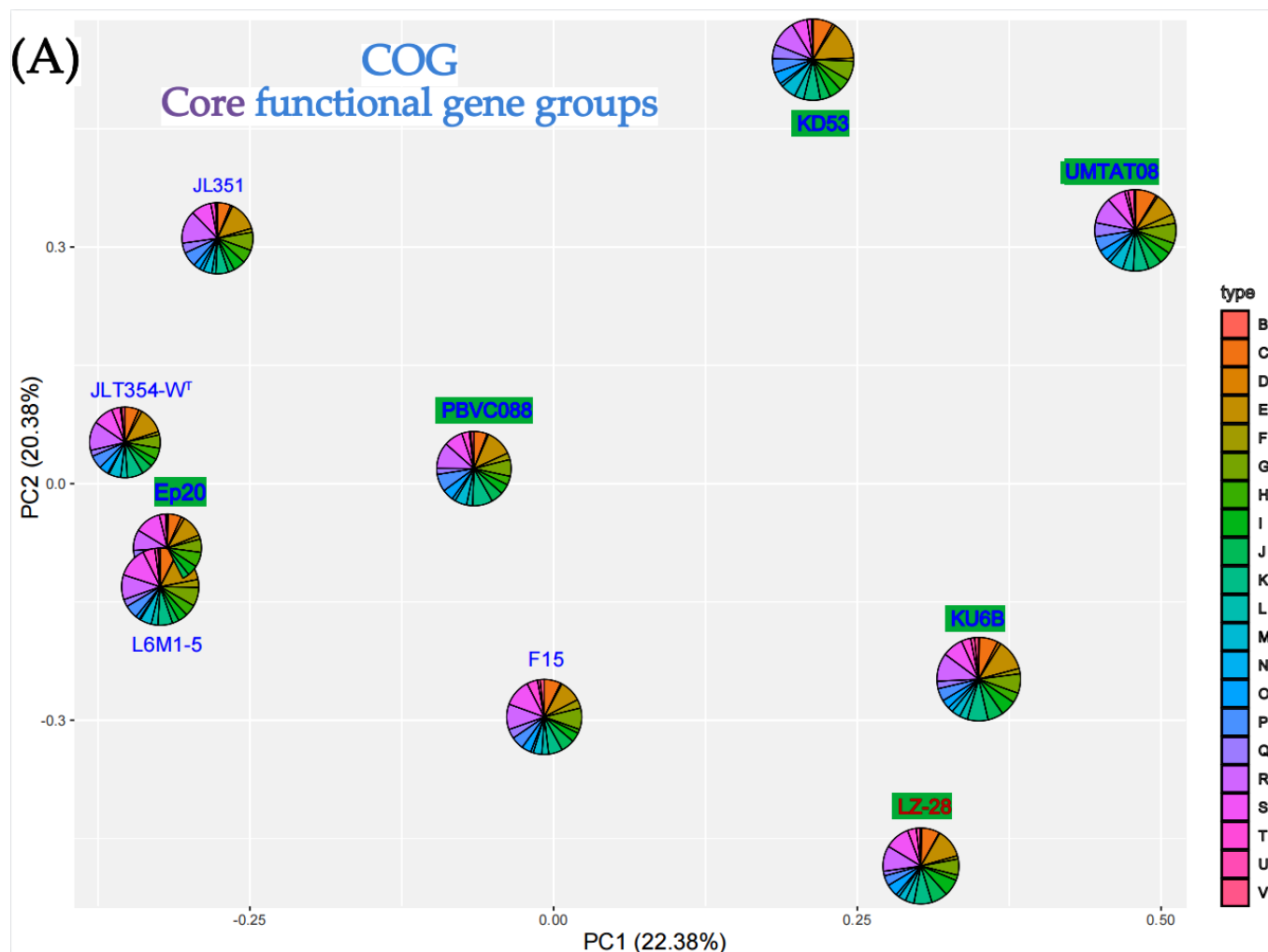
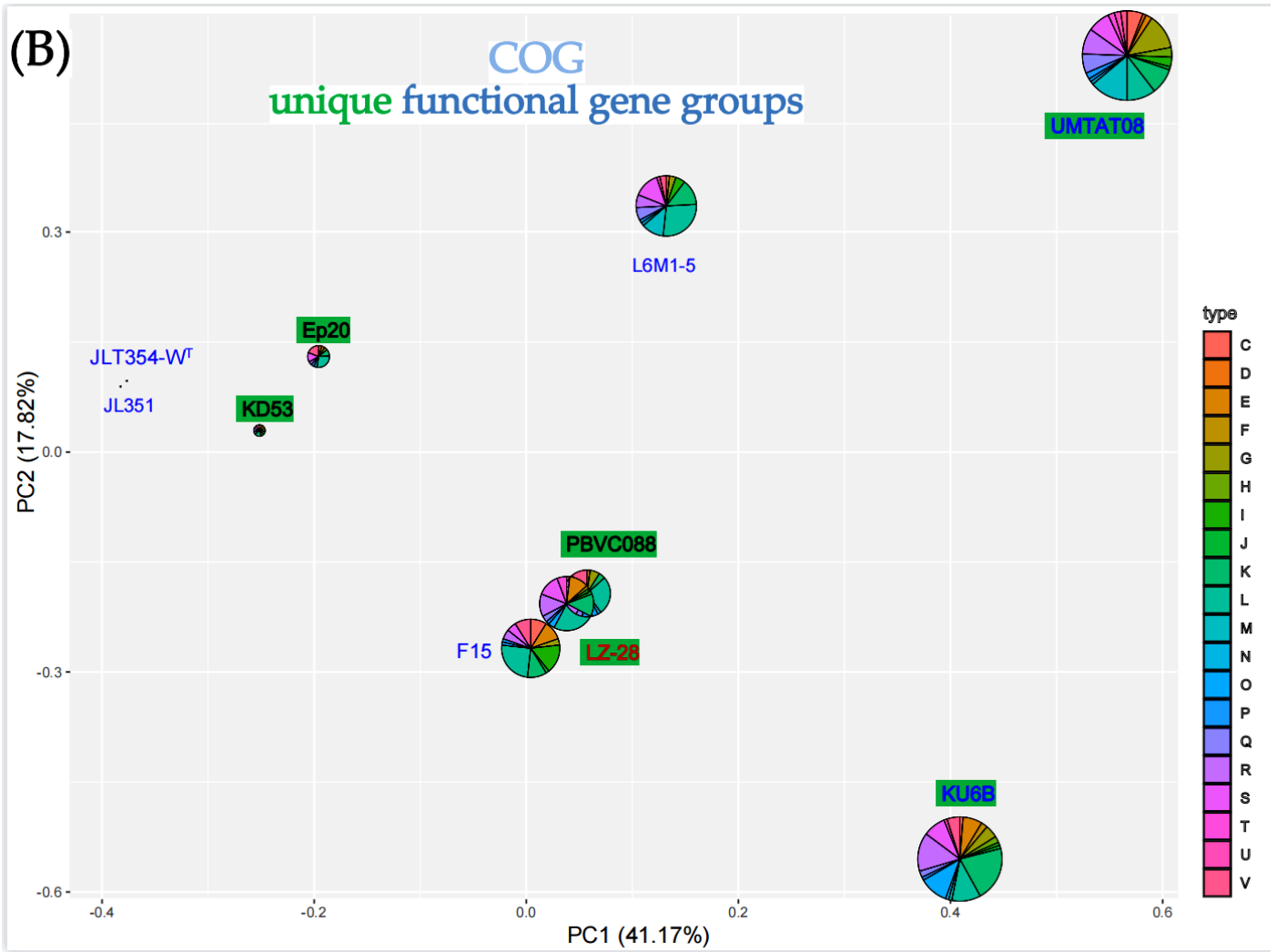


Figure S2. Comparison of the COG and KEGG categories among the ten selected *M. alba* strains. The functional genes of the core and unique groups distributed by COG database were shown in pane A and B, respectively. The pane C and D represented the distribution of the core genes and unique genes annotated by KEGG database, respectively. The sizes of the circles in each pane were proportional to the numbers of the different COG or KEGG categories. The strains isolated from marine algae were marked with green.

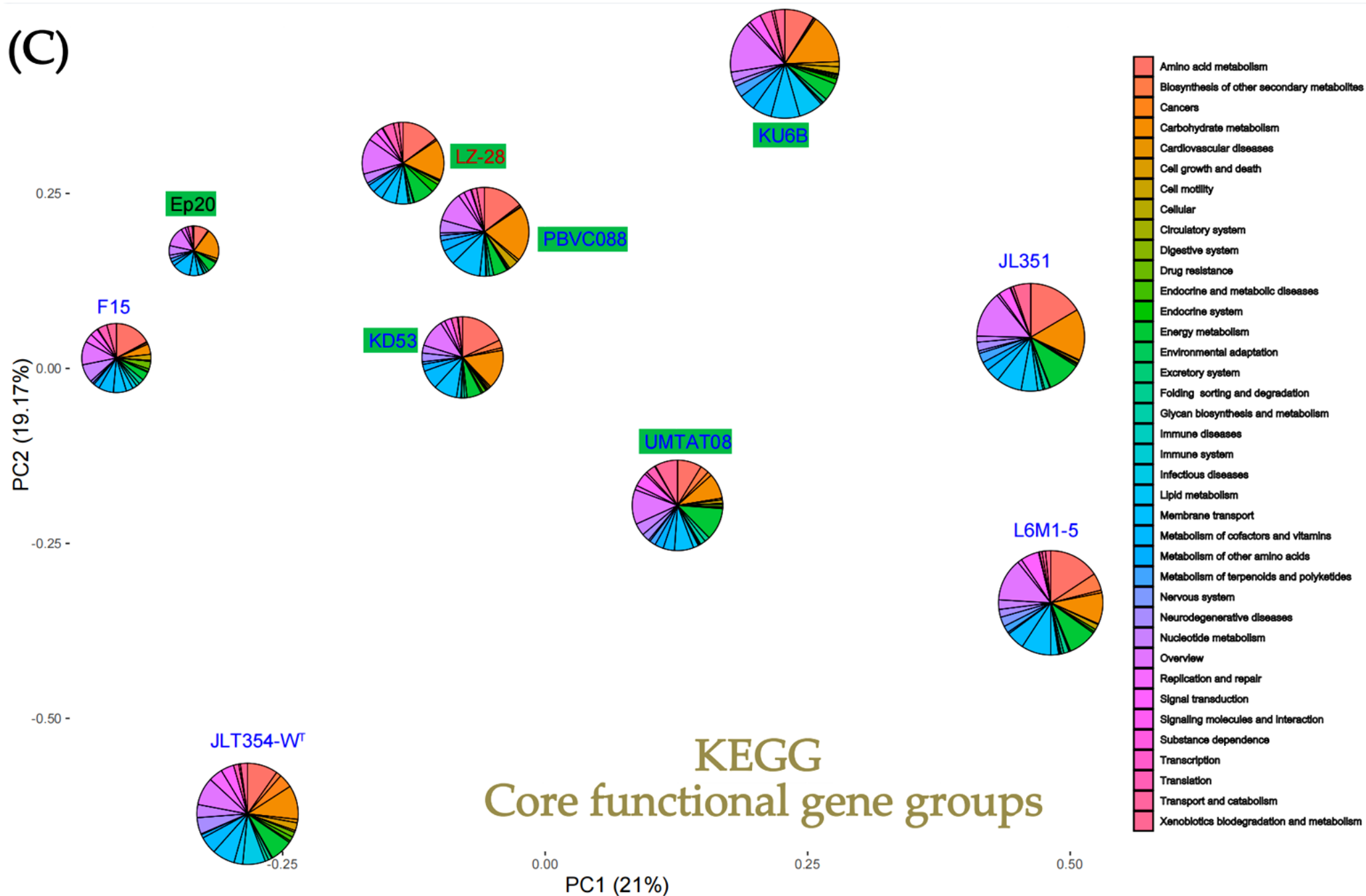


(B)

COG
unique functional gene groups



(C)



(D)

