

Diverse Cell Death Signature Based Subtypes Predict the Prognosis and Immune Characteristics Within Glioma

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Supplementary Description

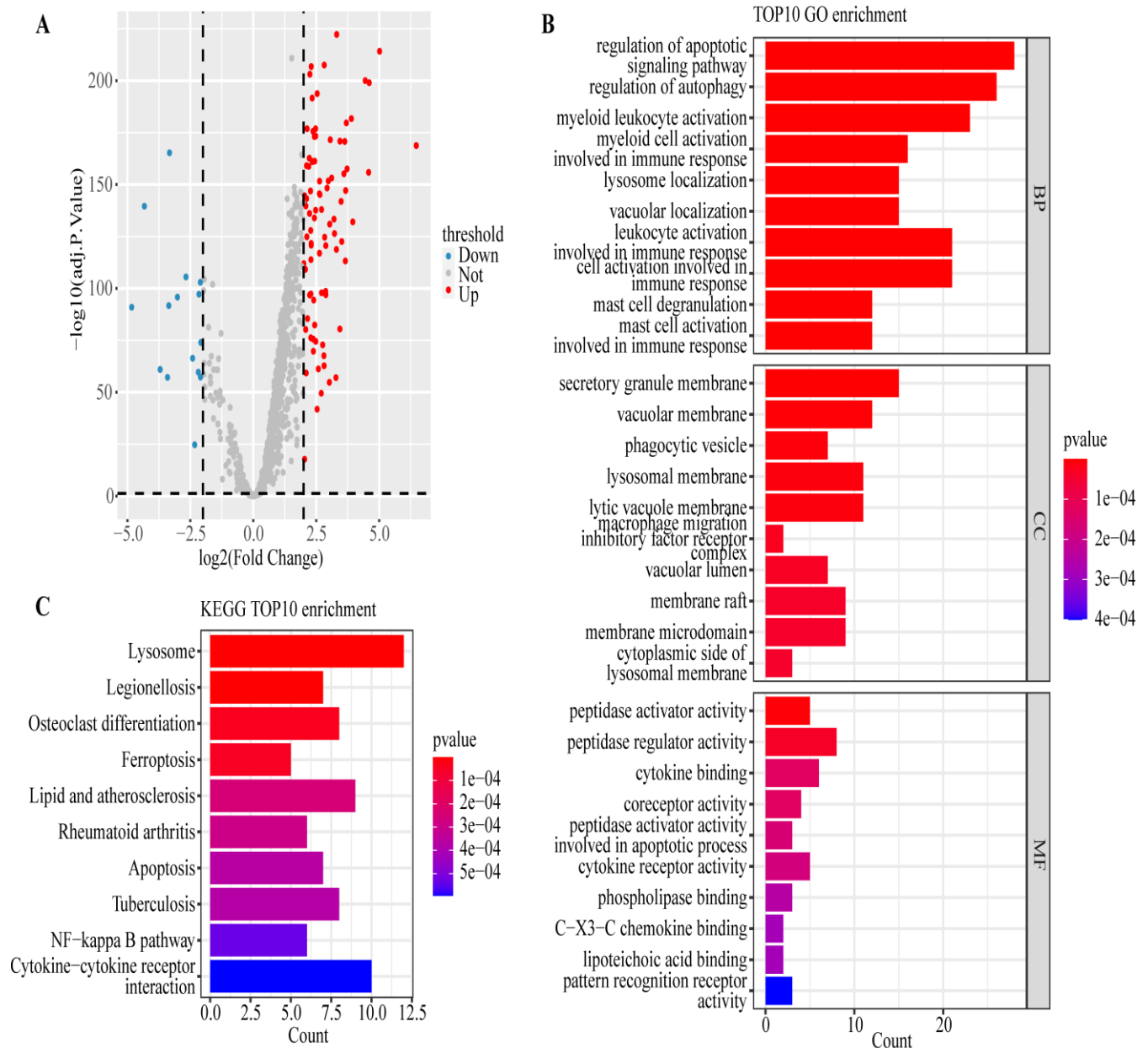


Figure S1: Differential expression analysis and functional enrichment analysis of PCD genes. (A) The volcano plot showing the PCD genes differentially expressed in glioma. (B-C) GO (B) and KEGG (C) enrichment analyses.

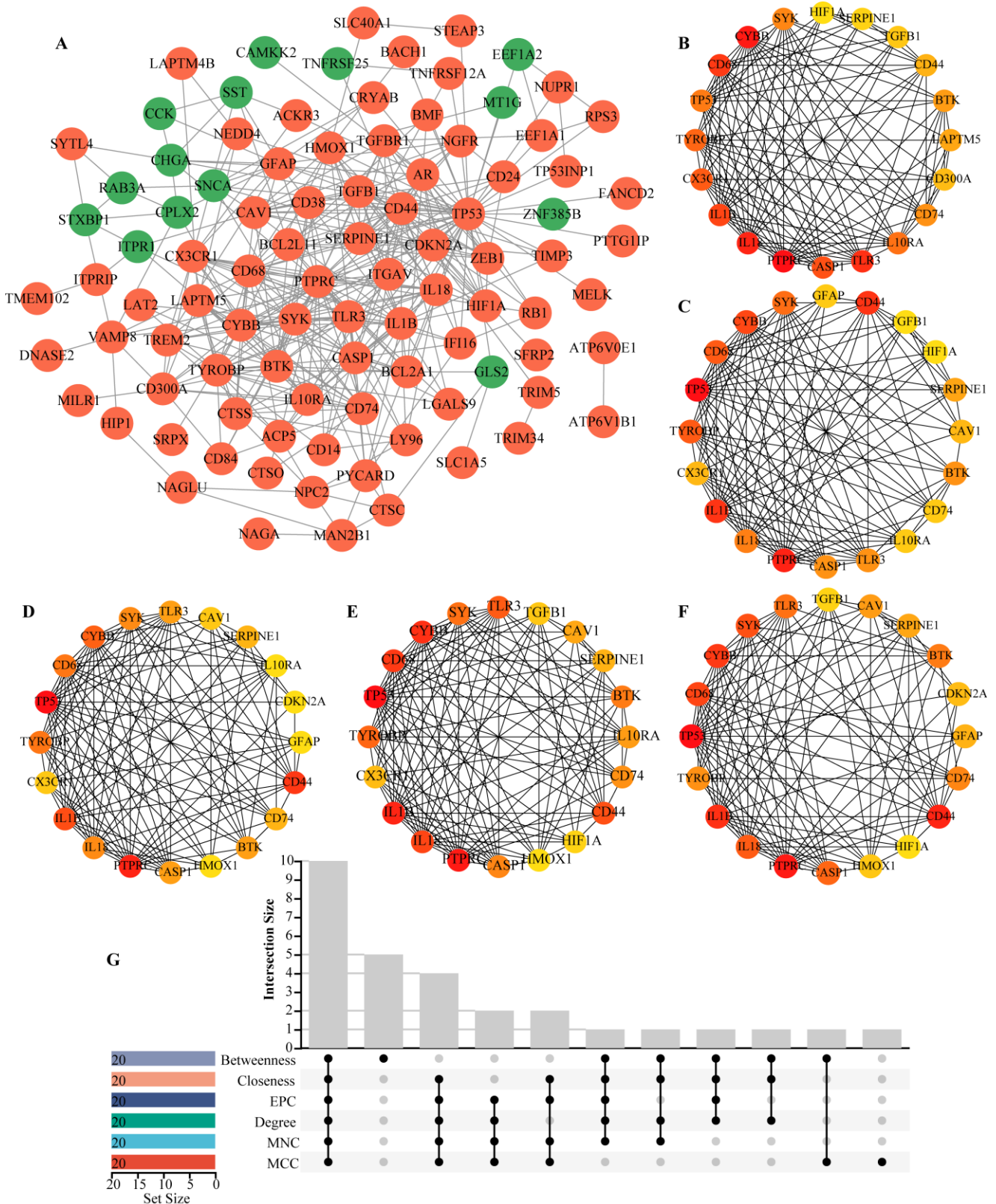


Figure S2: PPI network of differentially expressed PCD genes. (A) PPI network of differentially expressed PCD genes. (B-F) TOP20 gene networks by using MCC, MNC, Degree, EPC, and Closeness algorithms. (G) The upset plot of the key differentially expressed PCD genes.