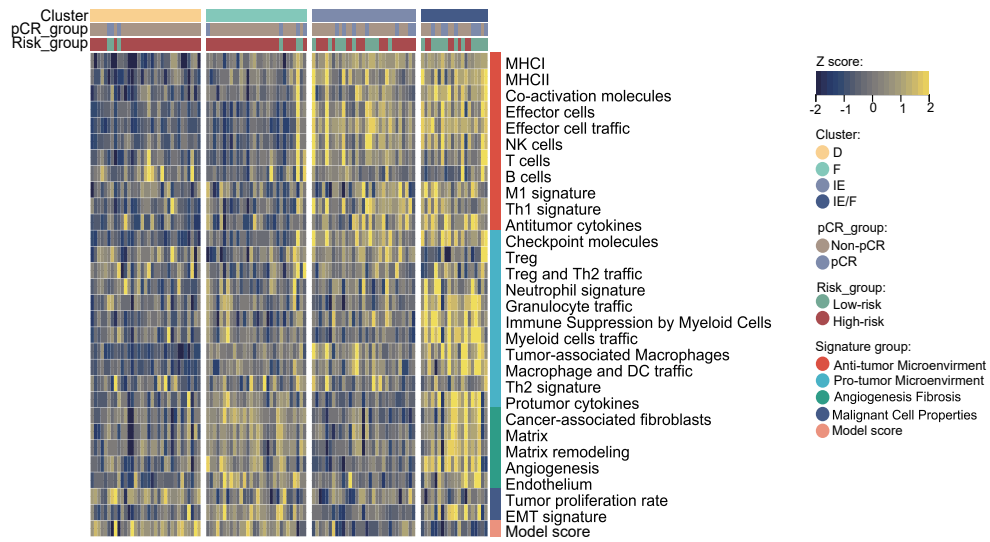
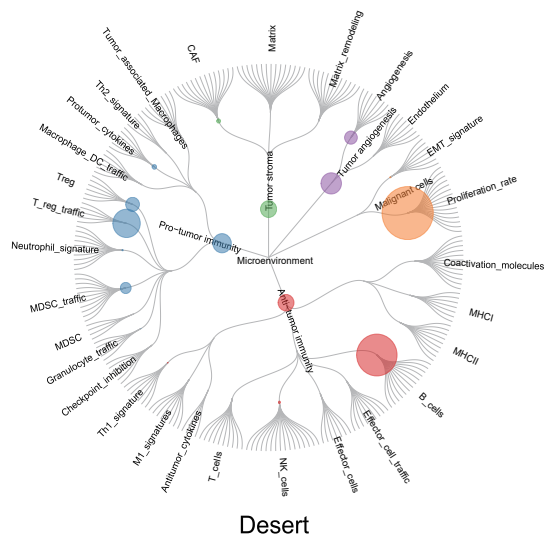


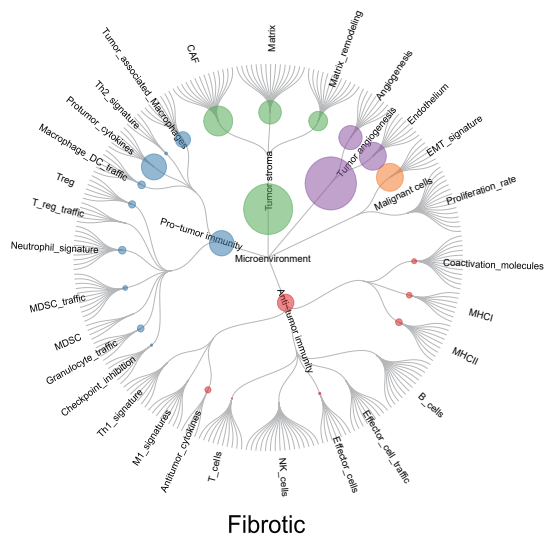
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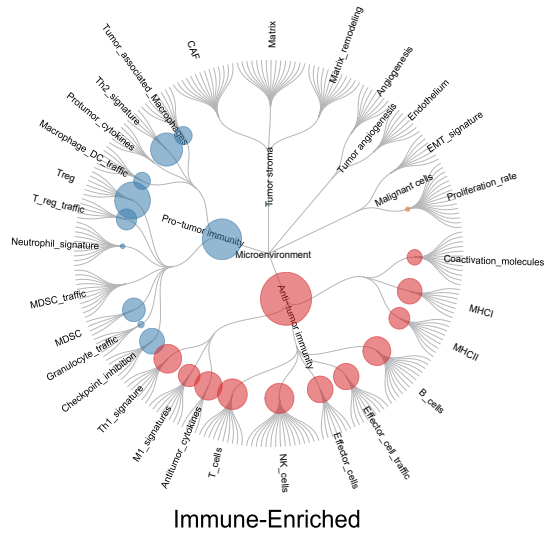
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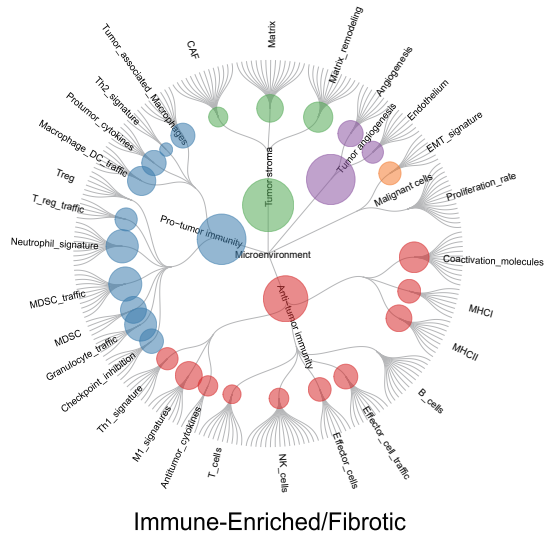
C



D



E



**Figure S6. TME quadricotype results for patients.** (A) Heatmap of 114 samples, including both cohorts 1 and 2, showcasing 29 gene expression signatures related to the TME. Based on the clustering analysis, the samples were categorized into four distinct subtypes: D (Desert), F (Fibrotic), IE (Immune-Enriched), and IE/F (Immune-Enriched/Fibrotic). Petal plots show molecular functional characteristics of the Desert (B), Fibrotic (C), Immune-Enriched (D), and Immune-Enriched/Fibrotic (E) subtypes.