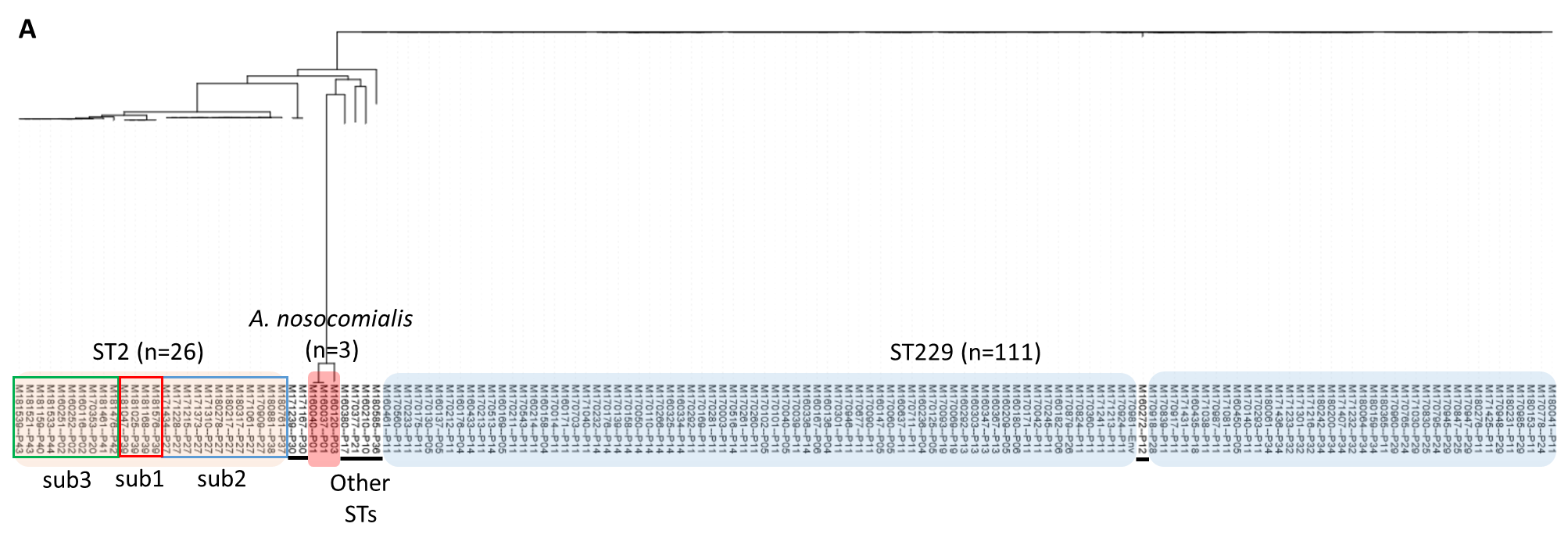
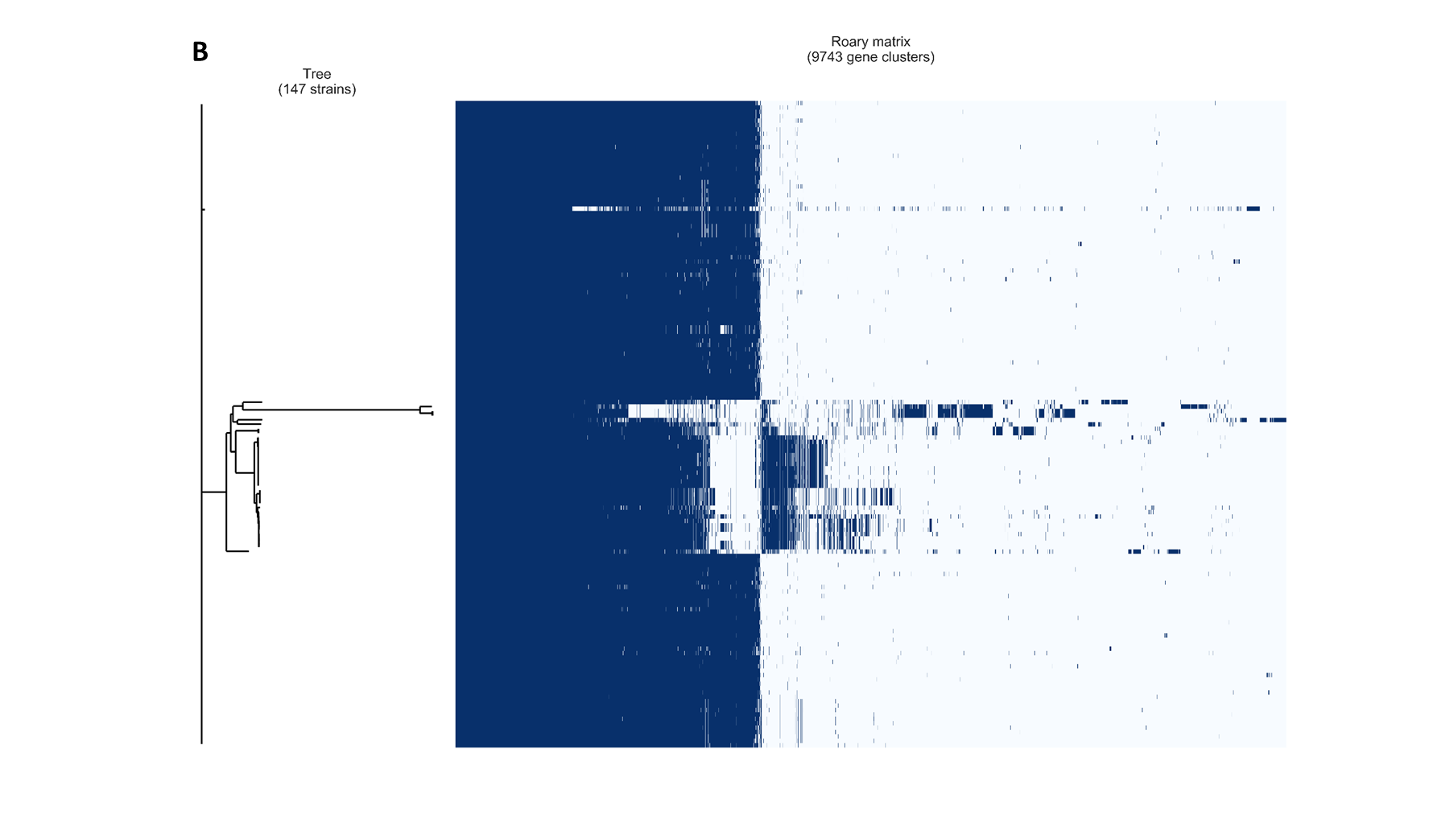
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**Supplementary Figure S1.** Roary core- and pan-genome analysis of 147 *Acinetobacter* clinical isolates. A) Phylogeny tree from Roary core- and pan-genome analysis. Four main clusters are identified: 3 *A. nosocomialis* isolates; 26 ST2 isolates; 111 ST229 isolates; and other *A.* *baumannii* isolates with other STs (underlined). Only three sub-clusters are identified in ST2 isolates: sub1, sub2 and sub3. B) Roary matrix plot of 9743 gene clusters identified from the 147 clinical isolates.