BMC infectious diseases supplementary

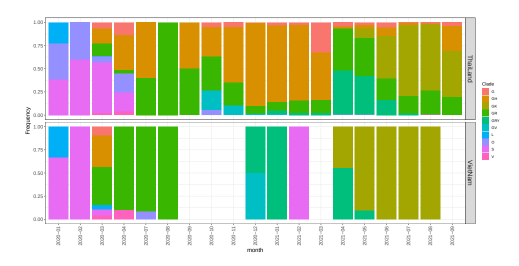
Tham Hoang

November 2021

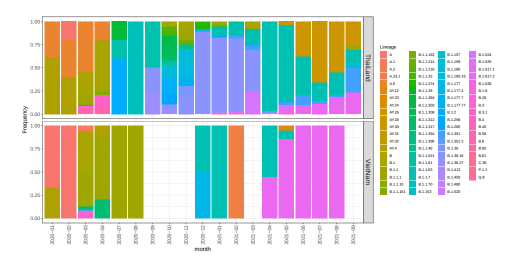
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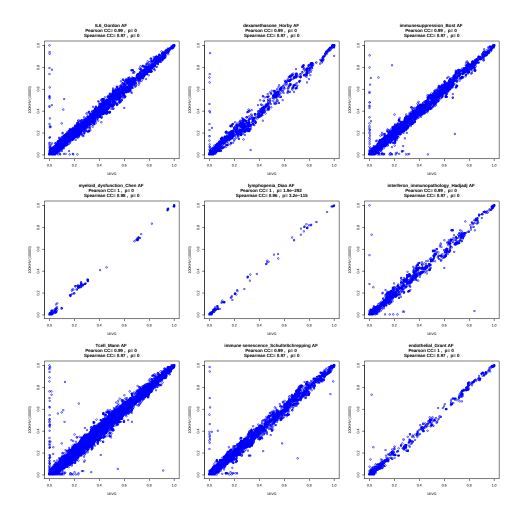
1 Supplementary Figures



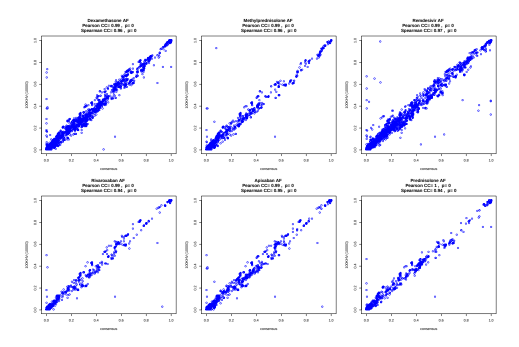
Supplementary Figure 1: Percentage of Clades by month in Vietnam and Thailand.



Supplementary Figure 2: Percentage by Lineage by month in Vietnam and Thailand.



Supplementary Figure 3: Scatter plots of allele frequency from datasets: one from WGS of Vietnamese people in the 1000 Genomes Project with high coverage, and the other from WGS of 1000 Vietnamese people [1] with nine immune gene sets associated with severity of COVID-19 in Vietnam.



Supplementary Figure 4: Scatter plots of allele frequency from datasets: one from WGS of Vietnamese people in the 1000 Genomes Project with high coverage, and the other from WGS of 1000 Vietnamese people [1] with 6 target gene sets of COVID-19 drugs used in Vietnam.

References

[1] 1kvg data portal. https://genome.vinbigdata.org/.