

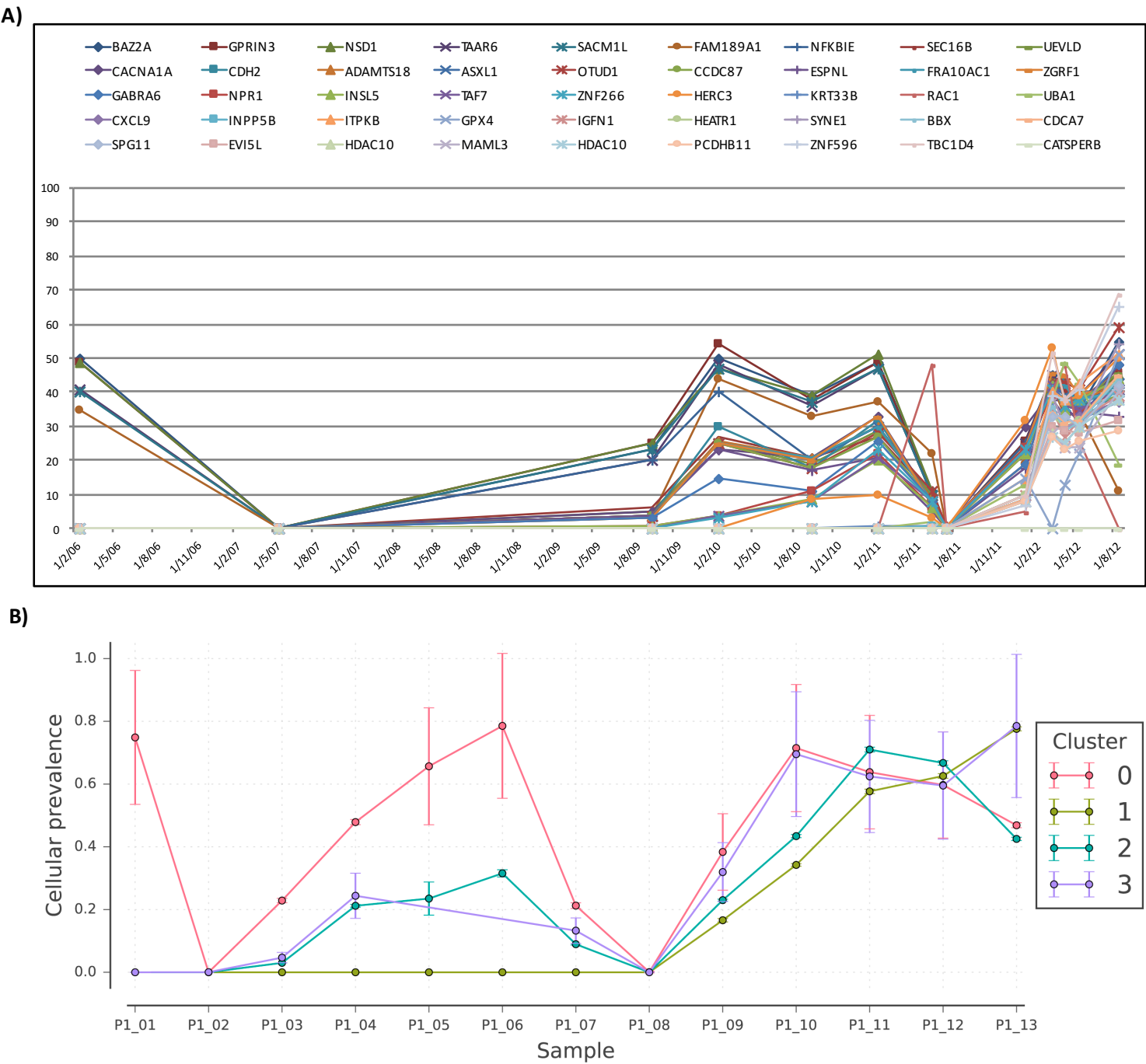
Supplementary Figures

Clonal dynamics monitoring during clinical evolution in chronic lymphocytic leukaemia

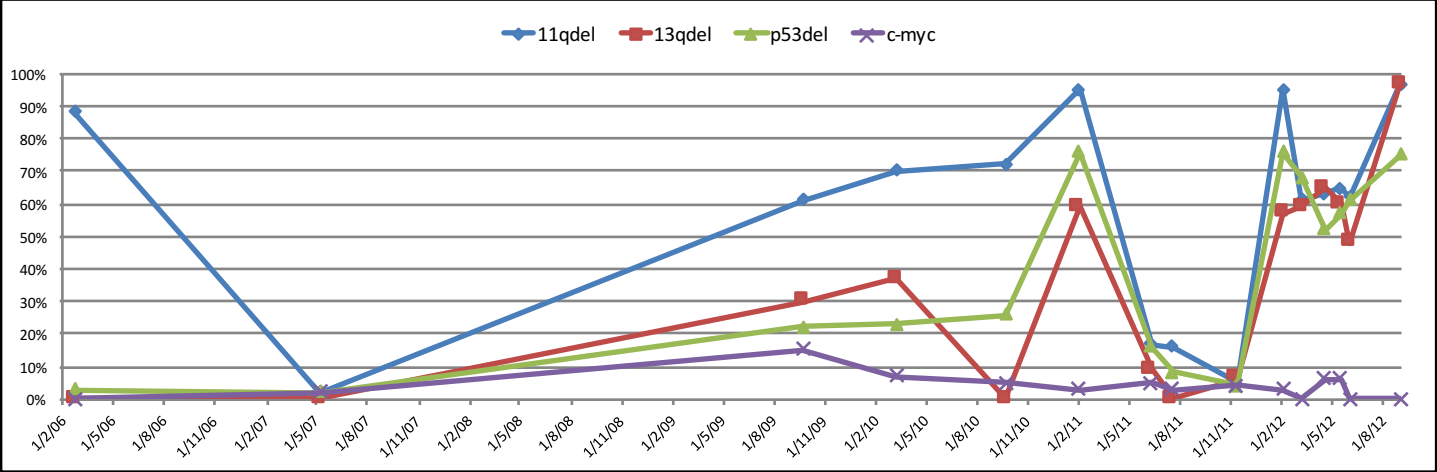
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Figure S1. Patient 1: A) VAF of the non-synonymous somatic mutations. B) Clonal composition inferred by PyClone during tumor evolution. C) percentage of cells with cytogenetics alterations detected by FISH during disease progression. D) Clonal composition inferred by PyClone of different affected organs



c)



d)

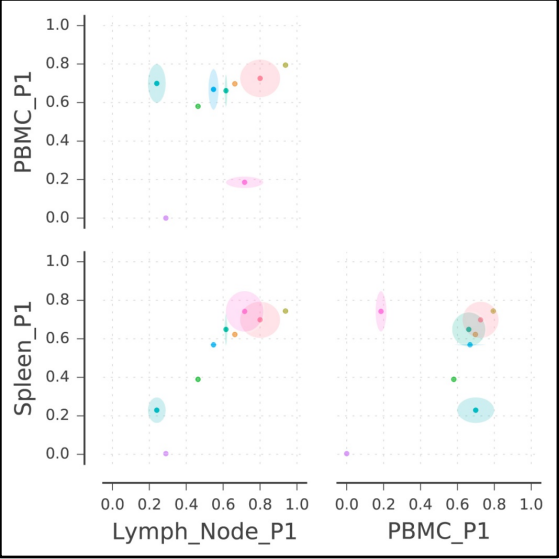
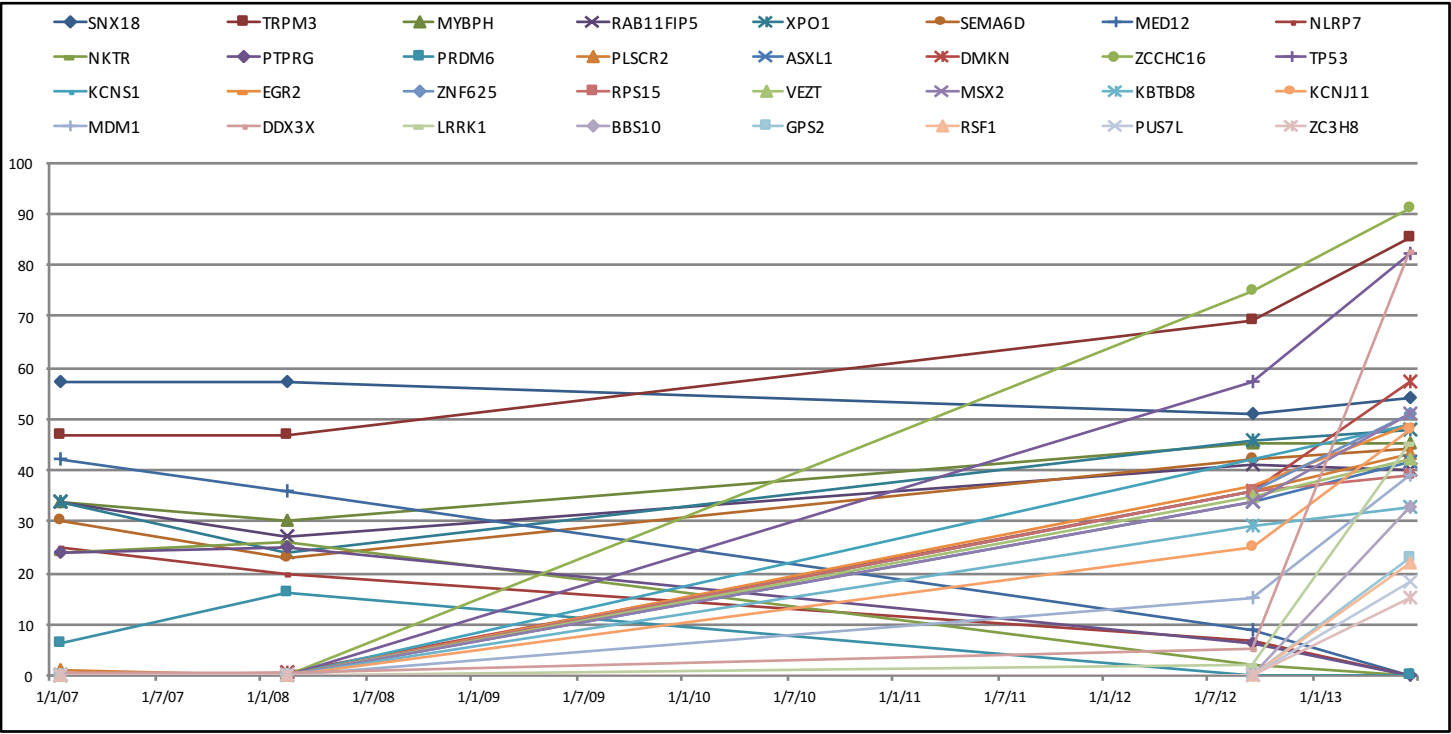
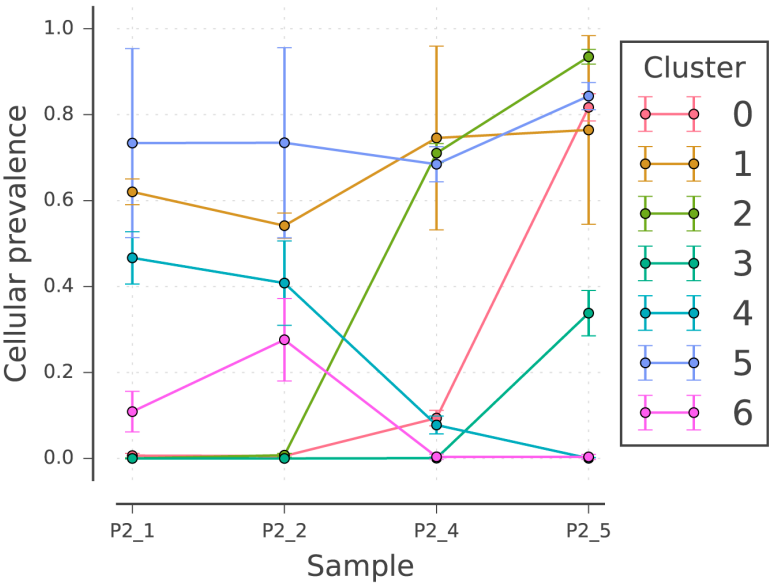


Figure S2. Patient 2: A) VAF of the non-synonymous somatic mutations. B) Clonal composition inferred by PyClone during tumor evolution. C) percentage of cells with cytogenetics alterations detected by FISH during disease progression.

A)



B)



c)

